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(54) Title: HUMAN BIKUNIN

(57) Abstract

The instant invention provides for proteins, polypeptides, nucleic acid sequences, constructs, expression vectors, host cells, pharmaceutical compositions of, and methods for using human placental bikunin, serine protease inhibitor domains, and fragments thereof.

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#### Title of the Invention: Human Bikunin

#### Field of the Invention

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The compositions of the invention relate to the field of proteins which inhibit serine protease activity. The invention also relates to the field of nucleic acid constructs, vectors and host cells for producing serine protease inhibiting proteins, pharmaceutical compositions containing the protein, and methods for their use.

# 10 Background of the Invention

#### Problem Addressed

Blood loss is a serious complication of major surgeries such as open heart surgery and other complicated procedures. Cardiac surgery patients account for a significant proportion of transfused donor blood. Blood transfusion carries risks of disease transmission and adverse reactions. In addition, donor blood is expensive and demands often exceed supply. Pharmacological methods for reducing blood loss and the resultant need for transfusion have been described (reviewed by Scott et al., Ann. Thorac. Surg. 50: 843-851, 1990).

### 20 Protein Serine Protease Inhibitor

Aprotinin, a bovine serine protease inhibitor of the Kunitz family is the active substance in the medicament Trasylol<sup>®</sup>. Aprotinin (Trasylol<sup>®</sup>) has been reported as being effective in reducing perioperative blood loss (Royston et al., Lancet ii: 1289-1291, 1987; Dietrich et al., Thorac. Cardiovasc. Surg. 37: 92-98, 1989; Fraedrich et al., Thorac. Cardiovasc. Surg. 37: 89-91, 1989); W. van Oeveren et al. (1987), Ann Thorac. Surg. 44, pp 640-645; Bistrup et al., (1988) Lancet I, 366-367), but adverse effects, including hypotension and flushing (Bohrer et al., Anesthesia 45: 853-854, 1990) and allergic reactions (Dietrich et al., Supra) have been reported. Use of aprotinin in patients previously exposed to it is not recommended (Dietrich et al., Supra). Trasylol<sup>®</sup> has also been used for the treatment of hyperfibrinolytic hemorrhages and traumatic hemorrhagic shock.

treatment of acute pancreatitis, various states of shock syndrome, hyperfibrinolytic hemorrhage and myocardial infarction (Trapnell et al., (1974) Brit J. Surg. 61: 177; J. McMichan et al., (1982) Circulatory Shock 9: 107. Auer et

al., (1979)Acta Neurochir. 49: 207; Sher (1977) Am J. Obstet. Gynecol. 129: 164; Schneider (1976), Artzneim.-Firsch. 26: 1606). It is generally thought that Trasylol® reduces blood loss *in vivo* through inhibition of kallikrein and plasmin. It has been found that aprotinin (3-58, Arg15, Ala17, Ser42) exhibits improved plasma kallikrein inhibitory potency as compared to native aprotinin itself (WO 89/10374).

#### Problems With Aprotinin

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Because aprotinin is of bovine origin, there is a finite risk of inducing anaphylaxis in human patients upon re-exposure to the drug. Thus, a human functional equivalent to aprotinin, by virtue of a lower risk of anaphylaxis, would be most useful and desirable to have.

Aprotinin is also nephrotoxic in rodents and dogs when administered repeatedly at high dose (Bayer, Trasylol®, Inhibitor of proteinase; Glasser et al., in "Verhandlungen der Deutschen Gesellschaft für Innere Medizin, 78. Kongress", Bergmann, Munchen, 1972 pp. 1612-1614). One hypothesis ascribes this effect to the accumulation of aprotinin in the negatively charged proximal tubules of the kidney, due to its high net positive charge (WO 93/14120).

Accordingly, an object of the present invention is to identify human proteins with functional activity similar to aprotinin. It was also an object of the instant invention to identify human proteins, that would be less charged, yet exhibit the same, highly similar, or improved protease specificities as found for aprotinin, especially with respect to the potency of plasmin and kallikrein inhibition. Such inhibitors could then be used repeatedly as medicaments in human patients with reduced risk of adverse immune response and reduced nephrotoxicity.

## Brief Summary of the Invention

The instant invention provides for a purified human serine protease inhibitor which can specifically inhibit kallikrein, that has been isolated from human placental tissue via affinity chromatography.

The instant invention provides a newly identified human protein herein called human placental bikunin that contains two serine protease inhibitor domains of the Kunitz class. In one particular embodiment, the instant invention embodies a protein having the amino acid sequence:

ADRERSIHDF CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN 50
YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF 100
NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE 150
ACMLRCFRQQ ENPPLPLGSK VVVLAGAVS 179
(SEQ ID NO: 1)

In a prefered embodiment the instant invention provides for native human placental bikunin protein having the amino acid sequence:

10 ADRERSIHDF CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN 50
YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF 100
NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE 150
ACMLRCFRQQ ENPPLPLGSK 170
(SEQ ID NO: 52)

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In one aspect, the biological activity of the protein of the instant invention is that it can bind to and substantially inhibit the biological activity of trypsin, human plasma and tissue kallikreins, human plasmin and Factor XIIa. In a preferred embodiment, the present invention provides for a native human placental bikunin protein, in glycosylated form. In a further embodiment the instant invention encompasses native human bikunin protein which has been formed such that it contains at least one cysteine-cysteine disulfide bond. In a preferred embodiment, the protein contains at least one intra-chain cysteinecysteine disulfide bond formed between a pair of cysteines selected from the group consisting of CYS11-CYS61, CYS20-CYS44, CYS36-CYS57, CYS106-CYS156, CYS115-CYS139, and CYS131-CYS152, wherein the cysteines are numbered according to the amino acid sequence of native human placental bikunin. One of ordinary skill will recognize that the protein of the instant invention may fold into the proper three-dimensional conformation, such that the biological activity of native human bikunin is maintained, where none, one or more, or all of the native intra-chain cysteine-cysteine disulfide bonds are present. In a most preferred embodiment, the protein of the instant invention is

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Active protein of the instant invention can be obtained by purification from human tissue, such as placenta, or via synthetic protein chemistry techniques, as illustrated by the Examples below. It is also understood that the

protein of the instant invention may be obtained using molecular biology techniques, where self-replicating vectors are capable of expressing the protein of the instant invention from transformed cells. Such protein can be made as non-secreted, or secreted forms from transformed cells. In order to facilitate secretion from transformed cells, to enhance the functional stability of the translated protein, or to aid folding of the bikunin protein, certain signal peptide sequences may be added to the NH2-terminal portion of the native human bikunin protein.

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In one embodiment, the instant invention thus provides for the native human bikunin protein with at least a portion of the native signal peptide sequence intact. Thus one embodiment of the invention provides for native human bikunin with at least part of the signal peptide, having the amino acid sequence:

15 AGSFLAWLGSLLLSGVLA -1
ADRERSIHDFCLVSKVVGRCRASMPRWWYNVTDGSCQLFVYGGCDGNSNN 50
YLTKEECLKKCATVTENATGDLATSRNAADSSVPSAPRRQDSEDHSSDMF 100
NYEEYCTANAVTGPCRASFPRWYFDVERNSCNNFIYGGCRGNKNSYRSEE 150
ACMLRCFRQQENPPLPLGSKVVVLAGAVS 179
20 (SEQ ID NO: 2)

In a prefered embodiment the instant invention provides for a native human placental bikunin protein with part of the leader sequence intact, having the amino acid sequence of SEQ ID NO: 52 with an intact leader segment having the amino acid sequence:

MAQLCGL RRSRAFLALL GSLLLSGVLA -1 (SEQ ID NO: 53)

In another embodiment, the instant invention provides for bikunin
protein with part of the leader sequence intact, having the amino acid sequence
of SEQ ID NO: 52 with the intact leader segment having the amino acid
sequence:

MLR AEADGVSRLL GSLLLSGVLA -1 (SEQ ID NO: 54)

In a preferred numbering system used herein the amino acid numbered +1 is assigned to the NH2-terminus of the amino acid sequence for native

human placental bikunin. One will readily recognize that functional protein fragments can be derived from native human placental bikunin, which will maintain at least part of the biological activity of native human placental bikunin, and act as serine protease inhibitors.

In one embodiment, the protein of the instant invention comprises a fragment of native human placental bikunin, which contains at least one functional Kunitz-like domain, having the amino acid sequence of native human placental bikunin amino acids 7-159, hereinafter called "bikunin (7-159)". Thus the instant invention embodies a protein having the amino acid sequence:

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IHDFCLVSKVVGRCRASMPRWWYNVTDGSCQLFVYGGCDGNSNN	50
YLTKEECLKKCATVTENATGDLATSRNAADSSVPSAPRRQDSEDHSSDMF	100
NYEEYCTANAVTGPCRASFPRWYFDVERNSCNNFIYGGCRGNKNSYRSEE	150
ACMLRCFRQ	159
(SEQ ID NO: 3)	

where the amino acid numbering corresponds to that of the amino acid sequence of native human placental bikunin. Another functional variant of this embodiment can be the fragment of native human placental bikunin, which contains at least one functional Kunitz-like domain, having the amino acid sequence of native human placental bikunin amino acids 11-156, bikunin (11-156)

25	CLVSKVVGRCRASMPRWWYNVTDGSCQLFVYGGCDGNSNN	50
	YLTKEECLKKCATVTENATGDLATSRNAADSSVPSAPRRQDSEDHSSDMF	100
	NYEEYCTANAVTGPCRASFPRWYFDVERNSCNNFIYGGCRGNKNSYRSEE	150
	ACMLRC	156
	(SEQ ID NO: 50).	3.0

One can recognize that the individual Kunitz-like domains are also fragments of the native placental bikunin. In particular, the instant invention

bikunin amino acids 7-64, hereinafter called "bikunin (7-64)". Thus in one embodiment the instant invention encompasses a protein which contains at least one Kunitz-like domain having the amino acid sequence:

IHDFCLVSKVVGRCRASMPRWWYNVTDGSCQLFVYGGCDGNSNN	50
YLTKEECLKKCATV	64
(SEQ ID NO: 4)	

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where the amino acid numbering corresponds to that of the amino acid sequence of native human placental bikunin. Another form of the protein of the instant invention can be a first Kunitz-like domain consisting of the amino acid sequence of native human placental bikunin amino acids 11-61, "bikunin (11-61)" having the amino acid sequence:

CLVSKVVGRCRASMPRWWYNVTDGSCQLFVYGGCDGNSNN 50
YLTKEECLKKC 61
(SEQ ID NO: 5)

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The instant invention also provides for a protein having the amino acid sequence of a Kunitz-like domain consisting of the amino acid sequence of native human placental bikunin amino acids 102-159, hereinafter called "bikunin (102-159)". Thus one embodiment the instant invention encompasses a protein which contains at least one Kunitz-like domain having the amino acid sequence:

YEEYCTANAVTGPCRASFPRWYFDVERNSCNNFIYGGCRGNKNSYRSEE	150
ACMLRCFRQ (SEQ ID NO: 6)	159

where the amino acid numbering corresponds to that of the amino acid sequence of native human placental bikunin. Another form of this domain can be a Kunitz-like domain consisting of the amino acid sequence of native human placental bikunin amino acids 106-156, "bikunin (106-156)" having the amino acid sequence:

CTANAVTGPCRASFPRWYFDVERNSCNNFIYGGCRGNKNSYRSEE	150
ACMLRC	156
(SEOIDNO:7)	

Thus one of ordinary skill will recognize that fragments of the native

human bikunin protein can be made which will retain at least some of the native protein biological activity. Such fragments can also be combined in different orientations or multiple combinations to provide for alternative proteins which retain some of, the same, or more biological activity of the native human bikunin protein.

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One will readily recognize that biologically active protein of the instant invention may comprise one or more of the instant Kunitz-like domains in combination with additional Kunitz-like domains from other sources. Biologically active protein of the instant invention may comprise one or more of the instant Kunitz-like domains in combination with additional protein domains from other sources with a variety of biological activities. The biological activity of the protein of the instant invention can be combined with that of other known protein or proteins to provide for multifunctional fusion proteins having predictable biological activity. Thus, in one embodiment, the instant invention encompasses a protein which contains at least one amino acid sequence segment the same as, or functionally equivalent to the amino acid sequence of either SEQ ID NO: 5 or SEQ ID NO: 7.

An open reading frame which terminates at an early stop codon can still code for a functional protein. The instant invention encompasses such alternative termination, and in one embodiment provides for a protein of the amino acid sequence:

ADRERSIHDFCLVSKVVGRCRASMPRWWYNVTDGSCQLFVYGGCDGNSNN	<b>5</b> 0
YLTKEECLKKCATVTENATGDLATSRNAADSSVPSAPRRQDS	92
(SEQIDNO:8)	

In one embodiment, the instant invention provides for substantially purified, or recombinantly produced native human bikunin protein with an intact segment of the leader sequence, and at least a portion of the native transmembrane region intact. Thus one embodiment of the invention provides for native human bikunin, with an intact leader sequence, and with at least part of the transmembrane domain (underlined), having an amino acid sequence

PCT/US97/03894 WO 97/33996

	1)EST		MILR A	EADGVSRLL G	SLLLSGVLA	- 1
	2) PCR		MAQLCGL RI	RSR <mark>AFLALL</mark> G	SLLLSGVLA	- 1
	1) EST 2) PCR 3) \(\lambda \cdot DNA\)		MAQLCGL RI	rs <b>raflal</b> l g	SLLLSGVLA	- 1
5	1) ADRERSIHDF	CLVSKVVGRC	RASMPRWWYN	VTDGSCQLFV	YGGCDGNSNN	5(
	2) ADRERSIHDF	CLVSKVVGRC	RASMPRWWYN	VTDGSCQLFV	YGGCDGNSNN	50
	3) ADRERSIHDF					
	1)YLTKEECLKK	CATVTENATG	DLATSRNAAD	SSVPSAPRRQ	DSEDHSSDMF	100
10	2)YLTKEECLKK	CATVTENATG	DLATSRNAAD	SSVPSAPRRQ	DSEDHSSDMF	100
	3) YLTKEECLKK	CATVTENATG	DLATSRNAAD	SSVPSAPRRQ	DSEDHSSDMF	100
	1) NYEEYCTANA	VTGPCRASFP	RWYFDVERNS	CNNFIYGGCR	GNKNSYRSEE	150
	2) NYEEYCTANA	VTGPCRASFP	RWYFDVERNS	CNNFIYGGCR	GNKNSYRSEE	150
15	3) NYEEYCTANA	VTGPCRASFP	RWYFDVERNS	CNNFIYGGCR	GNKNSYRSEE	150
	1) ACMLRCFRQQ	ENPPLPLGSK	VVVLAGLFVM	VLILFLGASM	VYLIRVARRN	200
	2) ACMLRCFRQQ	ENPPLPLGSK	VVVLAGLFVM	VLILFLGASM	VYLIRVARRN	200
20	3) ACMLRCFRQQ	ENPPLPLGSK	VVVLAGLFVM	VLILFLGASM	VYLIRVARRN	200
20	1)QERALRTVWS	SGDDKEQLVK	NTYVL		2	25
	2) QERALRTVWS	FGD			2	13
	3)QERALRTVWS		NTYVL			25

25 where sequence 1) is EST derived consensus SEQ ID NO: 45, 2) is PCR clone SEQ ID NO:47, and 3) is lambda cDNA clone SEQ ID NO:49. In a preferred embodiment a protein of the instant invention comprises one of the amino acid sequence of SEQ ID NO: 45, 47 or 49 wherein the protein has been cleaved in the region between the end of the last Kunitz domain and the transmembrane 30 region.

The instant invention also embodies the protein wherein the signal peptide is deleted. Thus the instant invention provides for a protein having the amino acid sequence of SEQ ID NO: 52 continuous with a transmembrane amino acid sequence:

35	EST	VVVLAGLFVM	VLILFLGASM	VYLIRVARRN	200
	EST	QERALRTVWS	SGDDKEQLVK	NTYVL	225
		(SEQ ID NO: 6	9)		
	a transmemb	rane amino acid	d sequence:		
	PCR	VVVLAGLFVM	VLILFLGASM	VYLIRVARRN	200
40	PCR	QERALRTVWS	FGD		213
		(SEQ ID NO: 6	<b>8</b> )		
	or a transme	mbrane amino a	acid sequence:		
	λcDNA	VVVLAGLFVM	VLILFLGASM	VYLIRVARRN	200
	λcDNA	QERALRTVWS	SGDDKEQLVK	NTYVL	225
45		(SEQ ID NO: 6	7).		

The protein amino acid sequences of the instant invention clearly teach one of the art the appropriate nucleic acid sequences which can be used in molecular biology techniques to produce the proteins of the instant invention. Thus, one embodiment of the instant invention provides for a nucleic acid sequence which encodes for a human bikunin having the consensus DNA sequence of Figure 3 (SEQ ID NO: 9), which translates into the amino acid sequence for native human placental bikunin sequence of Figure 3 (SEQ ID NO: 10). In another embodiment, the instant invention provides for a consensus nucleic acid sequence of Figure 4C (SEQ ID NO: 51) which encodes for an amino acid sequence of Figure 4D (SEQ ID NO: 45).

In a preferred embodiment, the instant invention provides for a nucleic acid sequence which encodes for native human placental bikunin having the DNA sequence of Figure 4F (SEQ ID NO: 48) which encodes for the protein sequence of SEQ ID NO: 49. In an another embodiment, the instant invention provides for a nucleic acid sequence of Figure 4E (SEQ ID NO: 46) which encodes for a protein sequence of SEQ ID NO: 47.

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One can easily recognize that certain allelic mutations, and conservative substitutions made in the nucleic acid sequence can be made which will still result in a protein amino acid sequence encompassed by the instant invention. One of skill in the art can recognize that certain natural allelic mutations of the protein of the instant invention, and conservative substitutions of amino acids in the protein of the instant invention will not significantly alter the biological activity of the protein, and are encompassed by the instant invention.

The instant invention also provides for pharmaceutical compositions containing human placental bikunin and fragments thereof that are useful for the reduction of perioperative blood loss in a patient undergoing surgery.

The present invention also provides methods for reducing perioperative blood loss in a patient undergoing surgery, wherein an effective amount of the disclosed human serine protease inhibitors of the present invention in a biologically compatible vehicle is administered to the patient.

The present invention also provides for variants of placental bikunin, and the specific Kunitz domains described above that contain amino acid abstractions that after the protease specificity reference sites of substitutions are indicated below as positions Aaa + through Aaa = in the amino acid sequence for native placental bikunin. Substitutions at Xaa 1 through Xaa 16 are also preferred for variants of bikunin (7-64), while substitutions at Xaa 17 through Xaa 32 are preferred for variants of bikunin (102-159).

Thus the present invention embodies protein having an amino acid sequence:

	Ala Asp Arg Glu Arg Ser Ile Xaa <sup>1</sup> Asp Phe	10
	Cys Leu Val Ser Lys Val Xaa <sup>2</sup> Gly Xaa <sup>3</sup> Cys	20
5	Xaa 4 Xaa 5 Xaa 6 Xaa 7 Xaa 8 Xaa 9 Trp Trp Tyr Asn	30
	Val Thr Asp Gly Ser Cys Gln Leu Phe Xaa 10	40
	Tyr Xaa 11 Gly Cys Xaa 12 Xaa 13 Xaa 14 Ser Asn Asn	50
	Tyr Xaa 15 Thr Lys Glu Glu Cys Leu Lys Lys	60
	Cys Ala Thr Xaa <sup>16</sup> Thr Glu Asn Ala Thr Gly	70
10	Asp Leu Ser Thr Ser Arg Asn Ala Ala Asp	80
	Ser Ser Val Pro Ser Ala Pro Arg Arg Gln	90
	Asp Ser Glu His Asp Ser Ser Asp Met Phe	100
	Asn Tyr Xaa 17 Glu Tyr Cys Thr Ala Asn Ala	110
	Val Xaa $^{18}$ Gly Xaa $^{19}$ Cys Xaa $^{20}$ Xaa $^{21}$ Xaa $^{22}$ Xaa $^{23}$ Xaa $^{24}$	120
15	Xaa 25 Trp Tyr Phe Asp Val Glu Arg Asn Ser	130
	Cys Asn Asn Phe Xaa <sup>26</sup> Tyr Xaa <sup>27</sup> Gly Cys Xaa <sup>28</sup>	140
	Xaa 29 Xaa 30 Lys Asn Ser Tyr Xaa 31 Ser Glu Glu	150
	Ala Cys Met Leu Arg Cys Phe Arg Xaa <sup>32</sup> Gln	160
	Glu Asn Pro Pro Leu Pro Leu Gly Ser Lys	170
20	Val Val Leu Ala Gly Ala Val Ser	179
	(SEQ ID NO: 11).	

where Xaa<sup>1</sup> - Xaa<sup>32</sup> each independently represents a naturally occurring amino acid residue except Cys, with the proviso that at least one of the amino acid residues Xaa<sup>1</sup>-Xaa<sup>32</sup> is different from the corresponding amino acid residue of the native sequence.

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In the present context, the term "naturally occurring amino acid residue" is intended to indicate any one of the 20 commonly occurring amino acids, i.e., Ala, Arg, Asn, Asp, Cys, Gln, Glu, Gly, His, Ile, Leu, Lys, Met, Phe, Pro, Ser, Thr, Trp, Tyr and Val.

By substituting one or more amino acids in one or more of the positions indicated above, it may be possible to change the inhibitor specificity profile of native placental bikunin or that of the individual Kunitz-like domains, bikunin(7-64) or bikunin (102-159) so that it preferentially inhibits other serine proteases such as, but not limited to, the enzymes of the complement cascade, TF/FVIIa, FXa, thrombin, neutrophil elastase, cathepsin G or proteinase-3.

Examples of preferred variants of placental bikunin include those

wherein Xaa 1 is an amino acid residue selected from the group consisting of His, Glu, Pro, Ala, Val or Lys, in particular wherein Xaa 1 is His or Pro; or wherein Xaa<sup>2</sup> is an amino acid residue selected from the group consisting of Val, Thr, Asp, Pro, Arg, Tyr, Glu, Ala, Lys, in particular wherein Xaa<sup>2</sup> is Val or Thr; or wherein Xaa<sup>3</sup> is an amino acid residue selected from the group consisting of Arg, Pro, Ile, Leu, Thr, in particular wherein Xaa3 is Arg or Pro; or wherein  $Xaa^4$  is an amino acid residue selected from the group consisting of Arg, Lys and Ser, Gln, in particular wherein  $Xaa^4$  is Arg or Lys; or wherein Xaa<sup>5</sup> is an amino acid residue selected from the group consisting of Ala, Gly, Asp, Thr, in particular wherein Xaa<sup>5</sup> is Ala; or wherein Xaa<sup>6</sup> is an amino acid 10 residue selected from the group consisting of Ser, Ile, Tyr, Asn, Leu, Val, Arg, Phe, in particular wherein Xaa6 is Ser or Arg; or wherein Xaa7 is an amino acid residue selected from the group consisting of Met, Phe, Ile, Glu, Leu, Thr and Val, in particular wherein Xaa<sup>7</sup> is Met or Ile; or wherein Xaa<sup>8</sup> is an amino acid residue selected from the group consisting of Pro, Lys, Thr, Gln, Asn, Leu, Ser 15 or Ile, in particular wherein Xaa8 is Pro or Ile; or wherein Xaa9 is an amino acid residue selected from the group consisting of Arg, Lys or Leu, in particular wherein Xaa<sup>9</sup> is Arg: or wherein Xaa<sup>10</sup> is an amino acid residue selected from the group consisting of Val, Ile, Lys, Ala, Pro, Phe, Trp, Gln, Leu and Thr, in particular wherein  $Xaa^{10}$  is Val; or wherein  $Xaa^{11}$  is an amino acid residue 20 selected from the group consisting of Gly, Ser and Thr, in particular wherein  $Xaa^{11}$  is Gly; or wherein  $Xaa^{12}$  is an amino acid residue selected from the group consisting of Asp, Arg, Glu, Leu, Gln, Gly, in particular wherein Xaa12 is Asp or Arg; or wherein Xaa 13 is an amino acid residue selected from the group consisting of Gly and Ala; or wherein Xaa 14 is an amino acid residue selected from the group consisting of Asn or Lys; or wherein Xaa 15 is an amino acid residue selected from the group consisting of Gly, Asp, Leu, Arg, Glu, Thr, Tyr, Val, and Lys, in particular wherein Xaa 15 is Leu or Lys; or wherein Xaa 16 is an amino acid residue selected from the group consisting of Val, Gln, Asp, Gly, Ile, Ala, Met, and Val, in particular wherein Xaa 16 is Val or Ala; or wherein Xaa 17 is an amino acid residue selected from the group consisting of His, Glu, Pro, Ala Tys and Val in particular wherein Yaa 17 is Glu or Proper wherein Yaa 19 amino acid residue selected from the group consisting of Mar This MST in Arg. Tyr, Glu. Ala or Lys, in particular wherein Xaa 18 is Thr; or wherein Xaa 19 is an amino acid residue selected from the group consisting of Arg, Pro, Ile, Leu or Thr, in particular wherein Xaa 19 is Pro; or wherein Xaa 20 is an amino acid residue selected from the group consisting of Arg, Lys, Gln and Ser, in

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particular wherein  $Xaa^{20}$  is Arg or Lys; or wherein  $Xaa^{21}$  is an amino acid residue selected from the group consisting of Ala, Asp, Thr or Gly; in particular wherein Xaa<sup>21</sup> is Ala; or wherein Xaa<sup>22</sup> is an amino acid residue selected from the group consisting of Ser, Ile, Tyr, Asn, Leu, Val, Arg or Phe, in particular wherein Xaa<sup>22</sup> is Ser or Arg; or wherein Xaa<sup>23</sup> is an amino acid residue selected from the group consisting of Met, Phe, Ile, Glu, Leu, Thr and Val, in particular wherein Xaa<sup>23</sup> is Phe or Ile; or wherein Xaa<sup>24</sup> is an amino acid residue selected from the group consisting of Pro, Lys, Thr, Asn, Leu, Gln, Ser or Ile, in particular wherein Xaa<sup>24</sup> is Pro or Ile; or wherein Xaa<sup>25</sup> is an amino acid residue selected from the group consisting of Arg, Lys or Leu, in particular wherein Xaa<sup>25</sup> is Arg: or wherein Xaa<sup>26</sup> is an amino acid residue selected from the group consisting of Val, Ile, Lys, Leu, Ala, Pro, Phe, Gln, Trp and Thr, in particular wherein Xaa<sup>26</sup> is Val or Ile; or wherein Xaa<sup>27</sup> is an amino acid residue selected from the group consisting of Gly, Ser and Thr, in particular wherein Xaa<sup>27</sup> is Gly; or wherein Xaa<sup>28</sup> is an amino acid residue selected from the group consisting of Asp, Arg, Glu, Leu, Gly or Gln, in particular wherein Xaa<sup>28</sup> is Arg; or wherein Xaa<sup>29</sup> is an amino acid residue selected from the group consisting of Gly and Ala; or wherein Xaa<sup>30</sup> is an amino acid residue selected from the group consisting of Asn or Lys; or wherein Xaa31 is an amino acid residue selected from the group consisting of Gly, Asp, Leu, Arg, Glu, Thr, Tyr, Val, and Lys, in particular wherein Xaa<sup>31</sup> is Arg or Lys; or wherein Xaa<sup>32</sup> is an amino acid residue selected from the group consisting of Val, Gln, Asp, Gly, Ile, Ala, Met, and Thr, in particular wherein Xaa<sup>32</sup> is Gln or Ala.

### 25 Description of the Drawings

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The invention will be better understood from a consideration of the following detailed description and claims, taken in conjunction with the drawings, in which:

Figure 1 depicts the nucleotide sequence of EST R35464 (SEQ ID NO: 12) and the translation of this DNA sequence (SEQ ID NO: 13) which yielded an open reading frame with some sequence similarity to aprotinin. The translation product contains 5 of the 6 cysteines in the correct spacing that is characteristic for Kunitz-like inhibitor domains (indicated in bold). The position normally occupied by the remaining cysteine (at codon 38) contained instead a phenylalanine (indicated by an asterisk).

Figure 2 depicts the nucleotide sequence of EST R74593 (SEQ ID NO: 14), and the translation of this DNA sequence (SEQ ID NO: 15) which yielded an

open reading frame with homology to the Kunitz class of serine protease inhibitor domains. The translation product contained 6 cysteines in the correct spacing that is characteristic for Kunitz-like inhibitor domains (indicated in bold). However, this reading frame sequence includes stop codons at codon 3 and 23.

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Figure 3 depicts a deduced nucleic acid sequence of human placental bikunin (SEQ ID NO: 9) labeled "consensus" and matched with the translated protein amino acid sequence labeled "translated" (SEQ ID NO: 10). Also as comparison are shown the nucleic acid sequence for ESTs H94519 (SEQ ID NO: 16), N39798 (SEQ ID NO: 17), R74593 (SEQ ID NO: 14) and R35464 (SEQ ID NO: 12). The underlined nucleotides in the consensus sequence correspond to the site of PCR primers described in the Examples. Underlined amino acids in the translated consensus sequence are residues whose identity have been confirmed by amino acid sequencing of purified native human placental bikunin.

15 Nucleotide and amino acid code are standard single letter code, "N" in the nucleic acid code indicates an unassigned nucleic acid, and "\*" indicates a stop codon in the amino acid sequence.

Figure 4A depicts the original overlay of a series of ESTs with some nucleic acid sequence homology to ESTs encoding human placental bikunin, or portions thereof. Shown for reference are the relative positions of bikunin (7-64) and bikunin (102-159), labeled KID1 and KID2 respectively.

Figure 4B depicts a subsequent more comprehensive EST overlay incorporating additional ESTs. Numbers on the upper X-axis refer to length in base pairs, starting at the first base from the most 5' EST sequence. The length of each bar is in proportion to the length in base pairs of the individual ESTs including gaps. The EST accession numbers are indicated to the right of their respective EST bars.

Figure 4C depicts the corresponding alignment of the oligonucleotide sequences of each of the overlapping ESTs shown schematically in Figure 4B. The upper sequence (SEQ ID NO: 51) labeled bikunin represents the consensus oligonucleotide sequence derived from the overlapping nucleotides at each position. The numbers refer to base-pair position within the EST map. The aligonucleotides in EST R74593 that are poid underlined (at map positions and 1005) are base insertions observed in R74593 that were consistently absent in each of the other overlapping ESTs.

Figure 4D depicts the amino acid translation of the consensus obligonucleotide sequence for bikunin depicted in Figure 4C (SEQ ID NO-45)

Figure 4E depicts the nucleotide sequence (SEQ ID NO: 46) and corresponding amino acid translation (SEQ ID NO: 47) of a placental bikunin encoding sequence that was derived from a human placental cDNA library by PCR-based amplification.

Figure 4F depicts the nucleotide sequence (SEQ ID NO: 48) and corresponding amino acid translation (SEQ ID NO: 49) of a native human placental bikunin encoding clone that was isolated from a human placental lambda cDNA library by colony hybridization.

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Figure 4G compares the alignment of the amino acid translated oligonucleotide sequences for placental bikunin obtained by EST overlay (SEQ ID NO: 45), PCR based cloning (SEQ ID NO: 47), and conventional lambda colony hybridization (SEQ ID NO: 49).

Figure 5 shows a graph of purification of human placental bikunin from placental tissue after Superdex 75 Gel-Filtration. The plot is an overlay of the protein elution profile as measured by OD 280 nm (solid line), activity of eluted protein in a trypsin inhibition assay (% inhibition shown by circles), and activity of eluted protein in a kallikrein inhibition assay (% inhibition shown by squares).

Figure 6 shows a graph which plots the purification of human placental bikunin from placental tissue using C18 Reverse-Phase Chromatography. The plot is an overlay of the protein elution profile as measured by OD 215 nm (solid line), activity of eluted protein in a trypsin inhibition assay (% inhibition shown by circles), and activity of eluted protein in a kallikrein inhibition assay (% inhibition shown by squares).

Figure 7 depicts a silver stained SDS-PAGE gel of highly purified placental bikunin (lane 2), and a series of molecular size marker proteins (lane 1) of the indicated sizes in kilodaltons. Migration was from top to bottom.

Figure 8 shows the amount of trypsin inhibitory activity present in the cell-free fermentation broth from the growth of yeast strains SC101 (panel 8A) or WHL341 (panel 8B) that were stably transformed with a plasmid (pS604) that directs the expression of placental bikunin (102-159).

Figure 9 shows both a silver stained SDS-PAGE (left panel) and a Western blot with anti-placental bikunin (102-159) pAb (right panel) of cell-free fermentation broth from the growth of yeast strain SC101 (recombinants 2.4 and 2.5) that was stably transformed with a plasmid directing the expression of either bovine aprotinin, or placental bikunin (102-159). Migration was from top to bottom.

Figure 10 is a photograph which shows a silver stained SDS-PAGE of highly purified placental bikunin (102-159) (lane 2) and a series of molecular

size marker proteins (lane 1) of the indicated sizes in Kilodaltons. Migration was from top to bottom.

Figure 11 is a photograph which shows the results of Northern blots of mRNA from various human tissues that was hybridized to a <sup>32</sup>P labeled cDNA probe encoding either placental bikunin (102-159) (panel 11A) or encoding placental bikunin (1-213) (panel 11B). Migration was from top to bottom. The numbers to the right of each blot refer to the size in kilobases of the adjacent RNA markers. The organs from which mRNA was derived is described under each lane of the blot.

Figure 12 depicts an immunoblot of placental derived placental bikunin with rabbit antiserum raised against either synthetic reduced placental bikunin (7-64) (panel A) or 102-159 (panel B). For each panel, contents were: molecular size markers (lanes 1); native placental bikunin isolated from human placenta (lanes 2); synthetic placental bikunin (7-64) (lanes 3) and synthetic placental bikunin (102-159) (lanes 4).

Tricine 10-20% SDS-PAGE gels were blotted and developed with protein A-purified primary polyclonal antibody (8 ug IgG in 20 ml 0.1% BSA/Tris-buffered saline (pH 7.5), followed by alkaline phosphatase-conjugated goat anti-rabbit secondary antibody. Migration was from top to bottom.

Figure 13 depicts a Coomassie Blue stained 10-20% Tricine SDS-PAGE gel of 3 micrograms of highly purified placental bikunin (1-170) derived from a baculovirus / Sf9 expression system (lane 2). Lane 1 contains molecular size markers. Migration was from top to bottom.

Figure 14 depicts a comparison of the effect of increasing concentrations of either Sf9-derived human placental bikunin (1-170) (filled circles), synthetic placental bikunin (102-159) (open circles), or aprotinin (open squares) on the activated partial thromboplastin time of human plasma. Clotting was initiated with CaCl<sub>2</sub>. The concentration of proteins are plotted versus the -fold prolongation in clotting time. The uninhibited clotting time was 30.8 seconds.

## 30 Detailed Description of the Invention

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The present invention encompasses a newly identified human protein herein called human placental bikunin that contains two serine protease auditor domains of the Kunitz class—he instant invention also encompassed marmaceutical compositions containing placental bikunin and tragments thereof that are useful for the reduction of perioperative blood loss in a patient undergoing surgery, or with major trauma.

The present invention also provides methods for reducing perioperative

blood loss in a patient undergoing surgery or due to major trauma, wherein an effective amount of the disclosed human serine protease inhibitors of the present invention, in a biologically compatible vehicle, is administered to the patient.

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A preferred application for placental bikunin, isolated domains, and other variants is for the reduction of blood loss resulting from trauma or surgery that has the potential for loss of large volumes of blood. These methods and compositions reduce or eliminate the need for whole donor blood or blood products, thereby reducing the risk of infection and other adverse side effects, as well as the cost of surgery. The methods are thus useful in reducing blood loss in normal patients, i.e., those not suffering from inborn or other preoperative deficiencies in coagulation factors. The reduction in blood loss is seen as a reduction in blood loss during surgery, as reduced post surgical drainage or both. Preferred surgical applications include but are not limited to use in thoracic and abdominal surgery, total and partial hip replacement surgeries and surgeries to treat a patient having an epithelial lesion of the eye. Preferred thoracic surgical procedures include but are not limited to aortocoronary bypass, excision of cardiac and aortic aneurysms, and surgery for esophageal varices, and coronary artery bypass surgery. Preferred abdominal surgeries include but are not limited to liver transplants, radical prostatectomy, surgery for diverticulitis of colon, tumor debulking, surgery on the abdominal aorta and surgery for duodenal ulcers, and repair of liver or spleen trauma. Preferred use for the treatment of trauma include but are not limited to the use in stabilization of severely injured patients at accident sites suffering from e.g., limb loss or major thoracic /abdominal wounds. In case of use for the reduction of blood loss resulting from surgery it is preferred to administer the placental bikunin, isolated domains, or other variant prior to and during surgery, whereas in case of use in trauma settings the placental bikunin variant, isolated domain or other variant is to be administered as rapidly as possible following injury, and should be contained on emergency vehicles traveling to the accident sites.

Factor XII (also known as Hageman Factor) is a serine protease that is found in the circulation in a zymogen form (80 kD) at approximately 29-40 µg/ml (see Pixley, et al. (1993) *Meth. in Enz.*, 222, 51-64) and is activated by tissue and plasma kallikrein. Once activated, it participates in the intrinsic pathway of blood coagulation which is activated when blood or plasma contacts a "foreign" or anionic surface. Once activated, Factor XIIa can then

cleave and activate a number of other plasma proteases including Factor XI, prekallikrein, and C1 of the complement system. Thus Factor XII may be involved in causing hypotensive reactions since activated kallikrein can cleave kiningen releasing bradykinin (see Colman, (1984) J. Clin. Invest., 73, 1249).

Sepsis is a disease that results from bacterial infection due to bacterial endotoxin or lipopolysaccharide (LPS). Exposure of Factor XII to LPS results in the activation of Factor XII. Patients with sepsis frequently have symptoms of intravascular coagulation which may also be due to activation of Factor XII by LPS. Septic shock can result from bacterial infection and is associated with fever, low systemic vascular resistance, and low arterial pressure. It is a common cause of death in intensive care units in the United States, where seventy five percent of the patients that die from septic shock have a persistent hypotension (see Parillo, et al. (1989) *Ann Rev. Med.*, 40, 469-485).

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Adult respiratory distress syndrome is characterized by pulmonary edema, hypoxemia, and decreased pulmonary compliance. The pathogenesis of the disease is currently unknown although the proteolytic pathways of coagulation and fibrinolysis are believed to play a role (see Carvalho, et al. (1988) *J. Lab Clin. Med.*, 112: 270-277).

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The proteins of the instant invention are also a novel human Kunitz type inhibitor of kallikrein, an activator of Factor XII. Thus another object of the current invention is to present a method for the prophylactic or therapeutic treatment of systemic inflammatory reactions such as septic shock, adult respiratory distress syndrome (ARDS), preeclampsia, multiple organ failure and disseminated intravascular coagulation (DIC). The therapeutic or prophylactic administration of the peptides of the instant invention would result in the modulation of these inflammatory conditions and be beneficial to the patient.

Plasmin plays an important role in extracellular matrix degradation and the activation of matrix-metallo protease (MMP) cascades. Collectively these proteases mediate migration of and tissue invasion by both endothelial cells during angiogenesis/neovascularization, and cancer cells during metastasis. Neovascularization is essential to support tumor growth and metastasis is a process which mediates the spreading of tumors and which is associated with extremely poor patient prognosis.

Several preclinical studies suggest that Kunitz like serine protease inhibitors with a protease specificity similar to aprotinin are useful as medicaments for cancer. For example, aprotinin reduced tumor growth and invasion, with increased tumor necrosis when administered to hamsters bearing a highly invasive fibrosarcoma or to mice bearing a similarly malignant mammary carcinoma (Latner et al., (1974), Br. J. Cancer 30: 60-67; Latner and Turner, (1976), Br. J. Cancer 33: 535-538). Furthermore, administration of 200,000 KIU of aprotinin i.p. to C57B1/6 Cr male mice on days 1 to 14 post-inoculation with Lewis lung carcinoma cells, reduced pulmonary metastases by 50% although had no effect on primary tumor mass (Giraldi et al., (1977) Eur. J. Cancer, 13: 1321-1323). Similarly, administration of 10,000 KIU i.p. on each of days 13-16 post-inoculation of C57BL/6J mice with Lewis tumor cells inhibited pulmonary metastases by 90% without affecting the primary tumor growth (Uetsuji et al., (1992), Jpn. J. Surg. 22: 429-442). In this same study, administration of plasmin or kallikrein with the same dosing schedule was argued to increase the number of pulmonary metastases. These results

prompted the authors to suggest that perioperative administration of aprotinin to cancer patients may reduce the likelihood of metastases. Black and Steger (1976, Eur. J. Pharmacol., 38: 313-319) found that aprotinin inhibited the growth of the transplanted rodent Murphy-Strum lymphosarcoma in rats and suggested that the effect involved the inhibition of the kinin-forming enzyme system. Twice daily i.p. injection of female ddY mice with 10,000 KIU of aprotinin for 7 weeks to mice each bearing a single autochtonous squamous cell carcinoma resulting from 3-methylcholanthrene treatment reduced the growth rate of the primary tumors by 90%. In some animals tumor regression was observed. While all vehicle treated animals had died within the seven weeks, all of the aprotinin treatment group remained alive. Reduced tumor growth was associated with hyperkeratosis (Ohkoshi, Gann (1980), 71: 246-250).

Clinically, a surgically cured group of 26 patients who received aprotinin i.v. exhibited a 70% survival two years post surgery with no recurrence of tumors whereas a placebo group of 26 patients at the same time exhibited only a 38% survival with a significant rate of tumor recurrence (Freeman et al. Br. Soc. Gastroenterol. (1980) supplement A: 902). In a case study (Guthrie et al., Br. J. Clin. Pract (1981) 35: 330-332), administration of bromocriptine plus aprotinin to a patient with advanced cancer of the cervix caused remission. Aprotinin was administered both as a 500,000 KTU i.p. bolus every eight hours concurrently with a continuous i.v. infusion of aprotinin at a rate of 200,000 KTU per 6 hr for a total of seven days once a month. Treatment was ended at the end of the fourth month due to the development of an allergic reaction to aprotinin. More recent evidence has further underscored a role of plasmin as a target for these effects of aprotinin on metastases.

The mechanism for these events could be related to the fact that aprotinin blocks the invasive potential of cancer cell lines (Liu G., et al., Int J. Cancer (1995), 60: 501-506). Furthermore, since the proteins of the instant invention are also potent inhibitors of plasmin and kallikrien, they are contemplated for use as anti-cancer agents. For example they are contemplated for use in blocking primary tumor growth by restricting neovascularization, amary humor invasion and in blocking metastasis through inhibition of tissue unitration. The compounds may be administered locally to numors a systemically. In a preferred mode of treatment, the protein would be administered perioperatively during tumor debulking to minimize the risk of metastasis. In such a regime, the blood sparing properties of the compound would be additionally advantageous in providing a clearer surgical field of

view. Another preferred mode of administration would be as a combination therapy with either MMP inhibitors or chemotherapy. An additional preferred mode of administration would be as a locally administered gene therapy designed to achieve selective expression of placental bikunin within the tumor cells, or their associated stroma and vascular beds.

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Preferred types of cancers targeted for therapy would be vasular-dependent solid tumors such as breast, colon, lung, prostate and ovarian carcinomas which exhibit a high metastatic potential, and those for which local delivery of a high concentration of the protein is feasible such as lung cancers through pulmonary delivery, colon carcinomas through hepatic delivery to liver metastasis, or skin cancers such as head and neck carcinomas or melanomas through subcutaneous delivery. Since the proteins of the present invention are of human origin they would be less likely to be associated with allergic or anaphylactic reactions of the kind observed by Guthrie et al., *supra*, upon reuse.

Additionally, the proteins of the present invention are contemplated for use in the reduction of thromboembolic complications associated with activation of the intrinsic pathway of coagulation. This would include prevention of pulmonary embolism in late stage cancer patients, a frequent cause of death (Donati MB., (1994), Haemostasis 24: 128-131).

Edema of the brain and spinal cord is a complication resulting from traumatic brain or spinal cord injury, stroke, cerebral ischemia, cerebral and sub-arachnoid hemhorrhage, surgery (including open heart surgery), infectious diseases such as encephalitis and meningitis, granulomatous diseases such as Sarcoid and focal or diffuse carcinomas, and is a contributor to the high level of morbidity and death following these events. Bradykinin is known to disrupt the blood brain barrier experimentally (Greenwood J., (1991), Neuroradiology, 33: 95-100; Whittle et al., (1992), Acta Neurochir., 115: 53-59), and infusion of bradykinin into the internal carotid artery induced brain edema in spontaneously hypertensive rats (SHR) subjected to common carotid artery occlusion (Kamiya, (1990), Nippon Ika Daigaku Zasshi. 57: 180-191). Elevated levels of bradykinin are found in extracellular fluids following trauma in a model involving traumatized rat spinal chord (Xu et al., (1991), J. Neurochem, 57: 975-980), and in plasma and tissue from rats with brain edema resulting from cerebral ischaemia (Kamiya et al., (1993), Stroke, 24: 571-575). Bradykinin is released from high molecular weight kininogen by serine proteases including kallikrein (Coleman (1984) J. Clin Invest., 73: 1249), and the serine protease inhibitor aprotinin was found to block the magnitude of brain edema resulting from

cerebralschemia in SHR rats (Kamiya, (1990), Nippon Ika Daigaku Zasshi. 57: 180-191; Kamiya et al., (1993), Stroke, 24: 571-575) and rabbits subjected to a cold lesion of the brain (Unterberg et al., (1986), J. Neurosurgery, 64: 269-276).

These observations indicate that brain edema results from local proteolytic release of kinins such as bradykinin from high molecular weight kininogen, followed by bradykinin-induced increases in blood brain barrier permeability. Accordingly, placental bikunin and fragments thereof are contemplated as medicaments for the prevention of edema in patients at risk for this condition, particularly those of high risk of mortality or brain injury. This would include head and spinal trauma patients, polytrauma patients, patients undergoing surgery of the brain or spinal cord and their associated vessels or other generalsurgeries including open-heart surgery, patients who have suffered from a stroke, cerebral or sub-arachnoid hemorrhage, infectious diseases of the brain, granulomatous disease of the brain or diffuse or focal carcinomas and tumors of the brain or any conditions such as multiple sclerosis involving breakdown of the blood brain barrier or patients suffering from any other inflammatory processes of the brain or spinal cord. Patients would receive an administration of placental bikunin either as an infusion or bolus injection, intravenously or intracranially. Additional doses of placental bikunin could be administered intermittently over the following one to three weeks. Dose levels would be designed to attain circulating concentrations in excess of those required to neutralize elevations in plasma levels or bradykinin and other vasoactive peptides formed through the action of serine proteases, and sufficient to reduce edema. Since the protein is of human origin, repeated administration in this course of therapy would not lead to development of an immune reaction to the protein. Placental bikunin and fragments thereof would be contemplated for monotherapy or prophylacsis as well as for use in combination with other medicaments such as neurotherapeutics and neuroprotectants.

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Recent evidence (Dela Cadena R. A. at al., (1995), FASEB J. 9: 446-452) has indicated that the contact activation pathway may contribute to the pathogenesis of arthritis and anemia, and that kallikrein inhibitors may be of therapeutic benefit. Accordingly, protease inhibitors of the present invention are contemplated according to their capacity to inhibit human kallikrein as neglicaments for the treatment of arthritis and anemia in numans

I reatment of male non-insulin diabetic (NIDDM) patients with aprotinial significantly improved total glucose uptake and decreased the metabolic clearance rate of insulin (Laurenti et al., (1996), Diabetic Medicine 13: 642-645). Accordingly, the human proteins of the present invention are contemplated for chronic use as

medicaments for the treatment of NIDDM.

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Daily treatment of patients at risk of preterm delivery with urinary trypsin inhibitor for two weeks significantly reduced recurrent uterine contractions (Kanayama et al., (1996), Eur J. Obstet. Gynecol. & Reprod. Biol. 67: 133-138). Accordingly, the human proteins of the present invention are contemplated for use in the prevention of preterm delivery.

Aprotinin has been shown to stimulate differentiation of mouse myoblasts in culture (Wells and Strickland, Development, (1994), 120: 3639-3647)), a process that is inhibited by TGFb. TGFb exists as an inactive pro-polypeptide which is activated by limited proteolysis. The mechanism of aprotinin action has been proposed to involve inhibition of proteases which process pro-TGFb to the mature active form. TGFb has been shown to be up-regulated in various fibrotic lesions and has long thought to be a potential target for anti-fibrotic therapies. In a rat model of pulmonary fibrosis for example, TGF-b concentrations paralleled the extent of bleomycin-induced inflammation. Furthermore, plasmin levels in the alveolar macrophage coincided with mature TGF-b levels, and the addition of the plasmin inhibitor a-2-antiplasmin abrogated the post translational activation of pro-TGFb by the macrophage (Khal et al., (1996), Am. J. Respir. Cell Mol. Biol. 15: 252-259.) The data suggest that plasmin contributes to the formation of active TGFb by alveolar macrophage, and that this process plays a pathologic role in the bleomycin-induced lung inflammation.

In light of these observations, placental bikunin and fragments thereof are contemplated as therapeutics for various fibrotic disorders, including pulmonary, hepatic, renal and dermal (scleroderma) fibrosis.

Aerosilized aprotinin was shown to protect >50% of mice infected with lethal doses of either influenza virus or paramyxovirus (Ovcharenko and Zhirnov, Antiviral Research, (1994), 23: 107-118). A suppression of the development of fatal hemorrhagic bronchopneumonia and a normalization of body weight gain were also noted with aerosilized aprotinin treatment. In light of these observations, placental bikunin and fragments thereof are contemplated as therapeutics for various respiratory related influenza-like diseases.

The human placental bikunin, isolated domains, and other variants of the invention are contemplated for use in the medical/therapeutic applications suggested for native aprotinin or aprotinin analogues with other inhibitory profiles, in particular those which necessitate usage of large doses. These would include diseases for which use of the human protein is indicated by virtue of its ability to inhibit human serine proteases such as trypsin, plasmin.

kallikrein, elastase, cathepsin G and proteinase-3, which include and are not limited to: acute pancreatitis (pancreatic elastase and trypsin), inflammation, thrombocytopenia, preservation of platelet function, organ preservation, wound healing, various forms of shock, including shock lung, endotoxin shock and post operative complications; disturbances of blood coagulation such as hyperfibrinolytic hemorrhage; acute and chronic inflammatory reactions, in particular for the therapy and prophylaxis of organ lesions, such as for example pancreatitis and radiation induced enteritis, complex-mediated inflammatory reactions such as immunovasculitis, glomerulonephritis and types of arthritis; collagenoses in particular rheumatoid arthritis; types of arthritis caused by metabolism-related deposits (for example gout); degeneration of the elastic constituents of the connective tissue parts of organs, such as in atherosclerosis (serum elastase) or pulmonary emphysema (neutrophil elastase); adult respiratory distress syndrome, inflammatory bowel disease, and psoriasis.

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A major unexpected finding was that the synthetic peptides encoding bikunin (7-64), and bikunin (102-159), could properly fold into the correct threedimensional conformation having active protease inhibitor bioactivity (Examples 2 and 1, respectively). Upon folding, each of these fragments of Bikurin underwent a reduction in mass of 6 mass units, consistent with the formation in each case, of three intrachain disulfide bonds between six cysteine residues of each fragment. Another surprising finding is that the synthetic peptides encoding bikunin (7-64), bikunin (102-159), and bikunin (1-170) are highly inhibitory of plasmin and both tissue and plasma kallikrein (Example 4, 3, and 10 respectively). Inhibition of plasmin and kallikrein by Trasylol® is thought to be involved in the mechanism by which Trasylol® reduces blood loss during open heart surgery. Our unexpected findings of the specificity of the Kunitz domains of the present invention make them suitable therapeutic agents for blood sparing during surgery or trauma where there is significant blood loss, or for any other condition where inhibition of plasmin and/or kallikrein would be beneficial.

Furthermore, we showed in this disclosure (Example 10) that placental bikunin (1.170) is a potent inhibitor of factor XIa and a moderate inhibitor of the intrinsic pathway coagulation, serving to interconvert inactive factor IX into active factor IXa. Thus, Placental Bikunin inhibits two key enzymes of the intrinsic pathway, kallikrein and factor XIa. Consistent with these observations, we also showed that placental bikunin (1-170) is a potent inhibitor of the activated partial

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thromboplastin time, which is a measure of the speed of coagulation driven by the intrinsic pathway. On the other hand, we showed that Placental bikunin (1-170) is an extremely weak inhibitor of the tissue factor VIIa complex, suggesting that it is not important in the regulation of the extrinsic coagulation cascade.

Based on these unexpected findings, placental bikunin is contemplated as a medicament for diseases in which activation of the intrinsic pathway of coagulation contributes significantly to the disease mechanism. Examples of such diseases would include post-traumatic shock and disseminated intravascular coagulation.

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A significant advantage of the Kunitz domains of the present invention is that they are human proteins, and also less positively charged than Trasylol® (Example 1), thereby reducing the risk of kidney damage on administration of large doses of the proteins. Being of human origin, the protein of the instant invention can thus be administered to human patients with significantly reduced risk of undesired immunological reactions as compared to administration of similar doses of Trasylol®. Furthermore, it was found that bikunin (102-159), bikunin (7-64), and bikunin (1-170) are significantly more potent inhibitors of plasma kallikrein than Trasylol® in vitro (Example 3, 4 and 10). Thus bikunin and fragments thereof are expected to be more effective in vivo at lowering blood loss in patients.

The amount of serine protease inhibitor administered should be sufficient to provide a supra normal plasma level. For the prophylactic reduction of bleeding during and following coronary aortic by-pass surgery (CABG), the proteins of the instant invention may be used in place of Trasylol® while taking into account the differences in potency. The use of Trasylol® is outlined in the Physicians Desk Reference, (1995), listing for Trasylol® supplement A. Briefly, with the patient in a supine position, the loading dose of placental bikunin, isolated domain or other variant is given slowly over about 20 to 30 minutes, after induction of anesthesia but prior to sternotomy. In general, a total dose of between about 2x106 KIU (kallikrein inhibitory units) and 8 X106 KTU will be used, depending on such factors as patient weight and the length of the surgery. Preferred loading doses are those that contain a total of 1 to 2 million kallikrein inhibitory units (KIU). When the loading dose is complete, it is followed by the constant infusion dose, which is continued until surgery is complete and the patient leaves the operating room. Preferred constant infusion doses are in the range of about 250,000 to 500,000 KIU per hour. The pump prime dose is added to the priming fluid of the

cardiopulmonary bypass circuit, by replacement of an aliquot of the priming fluid prior to the institution of the cardiopulmonary bypass. Preferred pump prime doses are those that contain a total of about one to two million KIU.

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The proteins of the instant invention are employed in pharmaceutical compositions formulated in the manner known to the art. Such compositions contain active ingredient(s) plus one or more pharmaceutically acceptable carriers, diluents, fillers, binders, and other excipients, depending on the administration mode and dosage form contemplated. Examples of therapeutically inert inorganic or organic carriers known to those skilled in the art include, but are not limited to, lactose, corn starch or derivatives thereof, talc, vegetable oils, waxes, fats, polyols such as polyethylene glycol, water, saccharose, alcohols, glycerin and the like. Various preservatives, emulsifiers, dispersants, flavorants, wetting agents, antioxidants, sweeteners, colorants, stabilizers, salts, buffers and the like can also be added, as required to assist in the stabilization of the formulation or to assist in increasing bioavailability of the active ingredient(s) or to yield a formulation of acceptable flavor or odor in the case of oral dosing. The inhibitor employed in such compositions may be in the form of the original compound itself, or optionally, in the form of a pharmaceutically acceptable salt. The proteins of the instant invention can be adminstered alone, or in various combinations, and in combination with other therapeutic compositions. The compositions so formulated are selected as needed for administration of the inhibitor by any suitable mode known to those skilled in the art.

Parenteral administration modes include intravenous (i.v.), subcutaneous (s.c.), intraperitoneal (i.p.), and intramuscular (i.m.) routes. Intravenous administration can be used to obtain acute regulation of peak plasma concentrations of the drug as might be needed. Alternatively, the drug can be administered at a desired rate continuously by i.v. catheter. Suitable vehicles include sterile, non-pyrogenic aqueous diluents, such as sterile water for injection, sterile-buffered solutions or sterile saline. The resulting composition is administered to the patient prior to and/or during surgery by intravenous injection or infusion.

mproved nair-life and targeting or the drug to bhagosomes such a neutrophils and macrophage involved in inflammation may be aided by entrapment of the drug in liposomes. It should be possible to improve the selectivity of liposomal targeting by incorporating into the outside of the liposomes ligands that bind to macromolecules specific to target organs/tissues such as the GI tract and lungs. Alternatively, i.m. or s.c. deposit injection with

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or without encapsulation of the drug into degradable microspheres (e.g., comprising poly-DL-lactide-co-glycolide) or protective formulations containing collagen can be used to obtain prolonged sustained drug release. For improved convenience of the dosage form it is possible to use an *i.p.* implanted reservoir and septum such as the percuseal system. Improved convenience and patient compliance may also be achieved by use of either injector pens (e.g., the Novo Pin or Q-pen) or needle-free jet injectors (e.g., from Bioject, Mediject or Becton Dickinson). Precisely controlled release can also be achieved using implantable pumps with delivery to the desired site via a cannula. Examples include the subcutaneously implanted osmotic pumps available from ALZA such as the ALZET osmotic pump.

Nasal delivery may be achieved by incorporating the drug into bioadhesive particulate carriers (<200 mm) such as those comprising cellulose, polyacrylate or polycarbophil, in conjunction with suitable absorption enhancers such as phospholipids or acylcarnitines. Commercially available systems include those developed by Dan Biosys and Scios Nova.

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Pulmonary delivery represents a nonparenteral mode of administration of the drug to the circulation. The lower airway epithelia are highly permeable to a wide range of proteins of molecular sizes up to about 20 kDa. Micron-sized dry powders containing the medicament in a suitable carrier such as mannitol, sucrose or lactose may be delivered to the distal alveolar surface using dry powder inhalers such as those of Inhale<sup>TM</sup>, Dura<sup>TM</sup>, Fisons (Spinhaler<sup>TM</sup>), and Glaxo (Rotahaler<sup>TM</sup>), or Astra (Turbohaler<sup>TM</sup>) propellant based metered dose inhalers. Solution formulations with or without liposomes may be delivered using ultrasonic nebulizers.

Oral delivery may be achieved by incorporating the drug into tablets, coated tablets, dragées, hard and soft gelatin capsules, solutions, emulsions, suspensions or enteric coated capsules designed to release the drug into the colon where digestive protease activity is low. Examples of the latter include the OROS-CT/Osmet<sup>TM</sup> system of ALZA, and the PULSINCAP<sup>TM</sup> system of Scherer Drug Delivery Systems. Other systems use azo-crosslinked polymers that are degraded by colon-specific bacterial azoreductases, or pH sensitive polyacrylate polymers that are activated by the rise in pH in the colon. The above systems may be used in conjunction with a wide range of available absorption enhancers. Rectal delivery may be achieved by incorporating the drug into suppositories.

In its preferred medicinal application, for reduction of perioperative

blood loss, the preferred mode of administration of the placental bikunin variants of the present invention is parenterally, preferably by i.v. route through a central line.

The amount of the pharmaceutical composition to be employed will depend on the recipient and the condition being treated. The requisite amount may be determined without undue experimentation by protocols known to those skilled in the art. Alternatively, the requisite amount may be calculated, based on a determination of the amount of target protease such as plasmin or kallikrein which must be inhibited in order to treat the condition. As the active materials contemplated in this invention are deemed to be nontoxic, treatment preferably involves administration of an excess of the optimally required amount of active agent.

Additionally, placental bikunin, isolated domains or other variants may be used to isolate natural substances such as its cognate proteases from human material using affinity based separation methods, as well as to elicit antibodies to the protease that can be further used to explore the tissue distribution and useful functions of Placental bikunin.

## Searching Human Sequence Data

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The existence of a distinct human protein homologous in function to 20 aprotinin, was deduced following a unique analysis of sequence entries to the expressed-sequence-tag data-base (hereafter termed dbEST) at the NCBI (National Center for Biological Information, Maryland). Using the TBlastN algorithm (BLAST, or Basic Local Alignment Search Tool uses the method of Altschul et a., (1990) J. Mol Biol 215: 403-410, to search for similarities between a 25 query sequence and all the sequences in a data-base, protein or nucleic acid in any combination), the data-base was examined for nucleotide sequences bearing homology to the sequence of bovine pre-pro-aprotinin, Trasylol®. This search of numerous clones was selectively narrowed to two particular clones which could possibly encode for a deduced amino acid sequence that would 30 correspond to a human protein homologous in function to aprotinin. The selected nucleic acid sequences were R35464 (SEQ ID NO: 12) and R74593 (SEQ

min diatella mier am .. he translated protein sequence in the longest open reading frame for reloans. (SEQ ID NO: 13) was missing one of the 6 cysteines that are critical for formation of the Kunitz-domain covalent structure, meaning that the nucleic acid sequence of R35464 could not yield a functional inhibitor. Similarly, the

longest translated open reading frame from clone R74593 (SEQ ID NO: 15) contained a stop codon 5' to the region encoding the Kunitz like sequence, meaning that this sequence, could not be translated to yield a functional secreted Kunitz domain. The significance of these sequences alone was unclear. It was possible that they represented a) the products of pseudogenes, b) regions of untranslated mRNA, or c) the products of viable mRNA which had been sequenced incorrectly.

### Discovery of Human Bikunin

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To specifically isolate and determine the actual human sequence, cDNA primers were designed to be capable of hybridizing to sequences located 5' and 3' to the segment of cDNA encoding our proposed Kunitz like sequences found within R35464 and R74593. The primers used to amplify a fragment encoding the Kunitz like sequence of R74593 were

CGAAGCTTCATCTCCGAAGCTCCAGACG (the 3'primer with a HindIII site; SEQ ID NO:33) and AGGATCTAGACAATAATTACCTGACCAAGGA (the 5'primer with an XbaI site; SEQ ID NO:34).

These primers were used to amplify by PCR (30 cycles) a 500 base pair product from a human placental cDNA library from Clontech (MATCHMAKER, Cat #HL4003AB, Clontech Laboratories, Palo Alto, CA), which was subcloned into Bluescript-SK+ and sequenced with the T3 primer with a Sequenase<sup>TM</sup> kit version 2.0. Surprisingly, the sequence of the fragment obtained using our primers was different from the sequence listed in the dbEST data base for clone R74593. In particular, our new sequence contained an additional guanosine base inserted 3' to the putative stop codon, but 5' to the segment encoding the Kunitz-like sequence (Figure 3). The insertion of an additional G shifted the stop codon out of the reading frame for the Kunitz-like domain (G at base pair 114 of the corrected sequence for R74593; Figure 3).

Subsequent query of the dbEST for sequences homologous to the Kunitz-like peptide sequence of R74593 yielded H94519 derived from human retina library and N39798. These sequences contained a Kunitz-like sequence that was almost identical to the Kunitz-like domain encoded in R35464 except that it contained all six of the characteristic cysteines. Overlay of each of the nucleotide sequences with that of R74593 (corrected by the insertion of G at b,p, 114) and R35464 was used to obtain a consensus nucleotide sequence for a partial human placental bikunin (SEQ ID NO: 9; Figure 3). The translated consensus sequence yielded an open reading frame extending from residue -18 to +179 (Figure 3;

full translation SEQ ID NO: 10) that contained two complete Kunitz-like domain sequences, within the region of amino acid residues 17-64 and 102-159 respectively.

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Further efforts attempted to obtain additional 5' sequence by querying dbEST with the sequence of R35464. Possible matches from such searches, that possessed additional 5' sequence were then in turn used to re-query the dbEST. In such an iterative fashion, a series of overlapping 5' sequences were identified which included clones H16866, T66058, R34808, R87894, N40851 and N39876 (Figure 4). Alignment of some of these sequences suggested the presence of a 5' ATG which might serve as a start site for synthesis of the consensus translated protein sequence. From this selected information, it was now possible to selectively screen for, and determine the nucleic acid and polypeptide sequences of a human protein with homologous function to aprotinin.

Re-interrogation of the dbEST revealed a number of new EST entries shown schematically in Figure 4B. Overlap with these additional ESTs allowed us to construct a much longer consensus oligonucleotide sequence (Figure 4C) that extended both 5' and 3' beyond the original oligonucleotide sequence depicted in Figure 3. In fact, the new sequence of total length 1.6 kilobases extended all the way to the 3' poly-A tail. The increased number of overlapping ESTs at each base-pair position along the sequence improved the level of confidence in certain regions such as the sequence overlapping with the 3' end of EST R74593 (Figure 3). Several overlapping ESTs in this region corroborated two critical base deletions relative to R74593 (located as bold underlined in Figure 4C, map positions 994 and 1005). Translation of the new consensus sequence (Figure 4D) in the bikunin encoding frame yielded a form of placental bikunin that was larger (248 amino acids) than the mature sequence (179 amino acids) encoded from the original consensus (SEQ ID NO: 1), and was terminated by an in-frame stop codon within the oligonucleotide consensus. The size increase was due to a frame shift in the 3' coding region resulting from removal of the two base insertions unique to EST R74593. The frame shift moved the stop codon of the original consensus (Figure 3) out of frame enabling read through into a new frame encoding the additional amino acid sequence.

risensus sequence (SEQ ID NO Therween residues + The rencoding m

Kunitz domains), but contained a new C-terminal extension exhibiting a putative 24 residue long transmembrane domain (underlined in Figure 4D) followed by a short 31 residue cytoplasmic domain. The precise sequence

around the initiator methionine and signal peptide was somewhat tentative due to considerable heterogeneity amongst the overlapping ESTs in this region.

Analysis of the protein sequence by Geneworks<sup>TM</sup>, highlighted asparagine residues at positions 30 and 67 as consensus sites for putative N-linked glycosylation. Asparagine 30 was not observed during N-terminal sequencing of the full length protein isolated from human placenta, consistent with it being glycosylated.

#### Cloning of Human Bikunin

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The existence of a human mRNA corresponding to the putative human bikunin nucleotide sequence inferred from the analysis of Figure 3, was confirmed as follows. The nucleic acid primer hybridizing 5' to the Kunitzencoding cDNA sequence of R35464 (b.p. 3-27 of consensus nucleotide sequence in Figure 3):

(a 5' primer derived from R35464 sequence with an XbaI site; SEQ ID NO: 35), and the nucleic acid primer hybridizing 3' to the Kunitz encoding sequence of R74593 (b.p. 680-700 of consensus nucleotide sequence in Figure 3), was used to PCR amplify, from a Clontech human placental library, a fragment of the size (ca. 670 b.p) expected from a cDNA consensus nucleotide sequence encoding the placental bikunin sequence of Figure 3 (Shown schematically in Figure 4A).

Using a 5' primer hybridizing to a sequence in R87894 that is 126 b.p 5' to the putative ATG start site discussed above, (shown schematically in Figure 4A at b.p. 110) plus the same 3' primer to R74593 as used above, it was possible to amplify a fragment from a Clontech human placental library of the expected size (approximately 872 b.p) predicted by EST overlay (Shown schematically in Figure 4).

Sequencing of the 872 b.p. fragment showed it to contain nucleotide segment corresponding to b.p. 110 to 218 of EST R87894 at its 5' end and b.p. 310 to 542 of the consensus sequence for placental bikunin inferred from the EST overlay analysis (of Figure 3), at its 3' end. This 3' nucleotide sequence contained all of the Kunitz-like domain encoded by placental bikunin (102-159).

To obtain a cDNA encoding the entire extracellular region of the protein, the following 5' PCR primer:

CACCTGATCGCGAGACCCC (SEQ ID NO: 36)
designed to hybridize to a sequence within EST R34808 was used with the same
3' primer to EST 74593 to amplify (30 cycles) an approximately 780 base-pair

cDNA product from the human placental cDNA library. This product was gel purified, and cloned into the TA vector (Invitrogen) for DNA sequencing by the dideoxy method (Sanger F., et al., (1977) Proc. Natl. Acad. Sci (USA), 74: 5463-5467) with the following primers:

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Vector Specific: GATTTAGGTGACACTATAG (SP6) (SEQ ID NO: 37)

TAATACGACTCACTATAGGG (T7) (SEQ ID NO: 38)

Gene Specific:

TTACCTGACCAAGGAGGAGTGC (SEQ ID NO: 39)

AATCCGCTGCATTCCTGCTGGTG (SEQ ID NO: 40)

CAGTCACTGGGCCTTGCCGT (SEQ ID NO: 41)

The resulting cDNA sequence is depicted in Figure 4E together with its translation product. At the nucleotide level, the sequence exhibited only minor differences from the consensus EST sequence (Figure 4D). Translation of the sequence yielded a coding sequence containing an in-frame initiator ATG site, signal peptide and mature placental bikunin sequence and transmembrane domain. The translated sequence of the PCR product was missing the last 12 amino acid residues from the cytoplasmic domain as a consequence of the choice of selection of the 3' primer for PCR amplification. This choice of 3' PCR primer (designed based on the sequence of R74593) was also responsible for the introduction of an artifactual S to F mutation at amino acid position 211 of the translated PCR-derived sequence. The signal peptide deduced from translation of the PCR fragment was somewhat different to that of the EST consensus.

To obtain a full length placental bikunin cDNA, the PCR derived product (Figure 4E) was gel purified and used to isolate a non-PCR based full length clone representing the bikunin sequence. The PCR derived cDNA sequence was labeled with <sup>32</sup>P-CTP by High Prime (Boehringer Mannheim) and used to probe a placental cDNA Library (Stratagene, Unizap<sup>TM</sup> λ library) using colony hybridization techniques. Approximately 2 X 106 phage plaques underwent 3 rounds of screening and plaque purification. Two clones were deemed full length (~1.5 kilobases) as determined by restriction enzyme analysis and based

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equencing or one or these clone by the dideoxy method vielded the oligonucleotide sequence depicted in Figure 4F. The translation product from this sequence yielded a protein with inframe initiator methionine, signal peptide and mature placental bikunin sequence. The mature placental bikunin

sequence was identical to the sequence of the mature protein derived by translation of the EST consensus although the signal peptide sequence lengths and sequences differed. Unlike the PCR derived product, the cDNA derived by colony hybridization contained the entire ectodomain, transmembrane domain, cytoplasmic domain and in-frame stop codon. In fact, the clone extended all the way to the poly-A tail. The initiator methionine was followed by a hydrophobic signal peptide which was identical to the signal peptide encoded in the PCR derived clone. Subsequently we expressed and purified a soluble fragment of placental bikunin, bikunin (1-170), from Sf9 cells (Example 9), and found it to be a functional protease inhibitor (Example 10). Furthermore, we isolated from human placenta a soluble fragment of placental bikunin which was also an active protease inhibitor (Example 7). Both the natural protein and the form of the protein expressed in Sf9 cells are probably glycosylated at the asparagine residue at position 30 based on the recoveries of PTH-amino acids during N-terminal sequencing (Examples 7 and 9).

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bikunin has the capacity to exist as a transmembrane protein on the surface of cells as well as a soluble protein. Other transmembrane proteins that contain Kunitz domains are known to undergo proteolytic processing to yield mixtures of soluble and membrane associated forms. These include two forms of the Amyloid Precursor Protein termed APP751 (Esch F., et al., (1990) Science, 248: 1122-1124) and APP 770 (Wang R., et al., (1991), J. Biol Chem, 266: 16960-16964).

Contact activation is a process which is activated by exposure of damaged vascular surfaces to components of the coagulation cascade.

Angiogenesis is a process that involves local activation of plasmin at endothelial surfaces. The specificity of placental bikunin and its putative capacity to anchor to cell surfaces, suggest that the physiologic functions of transmembranous placental bikunin may include regulation of contact activation and angiogenesis.

The amino acid sequences for placental bikunin (7-64), bikunin (102-159), and full length placental bikunin (Figure 4F) were searched against the PIR (Vers. 46.0) and PatchX (Vers. 46.0) protein databases as well as the GeneSeq (Vers. 20.0) protein database of patented sequences using the Genetics Computer Group program FastA. Using the Genetics Computer Group program TFastA (Pearson and Lipman, 1988, Proc. Natl. Acad. Sci. USA 85: 2444-2448), these same protein sequences were searched versus the six-frame translations of the GenBank (Vers. 92.0 with updates to 1/26/96) and EMBL

(modified Vers. 45.0) nucleotide databases as well as the GeneSeq (Vers. 20.0) nucleotide database of patented sequences. The EST and STS subsets of GenBank and EMBL were not included in this set of searches. The best matches resulting from these searches contained sequences which were only about 50% identical over their full length to the 58-amino acid protein sequence derived from our analysis of clones R74593 and R35464.

#### Isolation of Human Bikunin

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As mentioned above, synthetic peptides corresponding to bikunin (7-64) and bikunin (102-159) as determined from the translated consensus sequence for bikunin (Figure 3), could be refolded (Examples 2 and 1, respectively) to yield active kallikrein inhibitor protein (Example 4 and 3, respectively). We exploited this unexpected property to devise a purification scheme to isolate native placental bikunin from human tissue.

Using a purification scheme which employed kallikrein-sepharose affinity chromatography as a first step, highly purified native potent kallikrein inhibitor was isolated. The isolated native human bikunin had an identical N-terminus (sequenced for 50 amino acid residues) as the sequence predicted by the translation of the consensus nucleic acid sequence (Figure 3) amino acid residues +1 to +50 (Example 7). This confirmed for the first time the existence of a novel native kallikrein inhibitor isolated from human placenta.

Known Kunitz-like domains are listed below. Residues believed to be making contact with target proteases are highlighted as of special interest (bold/underlined). These particular residues are named positions Xaa1-16 for specific reference as shown by label Xaa below:

	<b>X88</b> 1 2	3 456789		1 1 111 0 1 234	1 5	1 6
	1) I HDFCLVSKVV	GRCRASKPRW	WYNVTDGSCQ	LFVYGOC DON	SNNY LTKEEC	LKKCATV
5	2) Y BEYCTANAVT	GPCRASFPRW	YFDVERNSCN	NF IYGOC RCN	KNSY RSEEAC	MLRCFRQ
	3) - HSFCAFKADD	G <b>PCKAING</b> RF	FFNIFTRQCE	ef lygoc bon	QNRF ESLEEC	KKMCTR D
	4) - PDFCFLEED P	GICRGYITRY	FYNNQTKQCE	RF KYCCC LCH	MANF ETLEEC	KNICEDG
	5) - PSWCLTPADR	GLCRAFTERF	YYNSVIGKCR	PF KY SCC COM	<b>ENNFT</b> SKQEC	LRACKKG
	6) - AEICLLPLDY	GPCRALLLRY	YYRYRTQSCR	QF LYGCC BOM	ANNF YTWEAC	DDACWRI
10	7) - PSFCYSPKDE	GLCSANVTRY	YFNPRYRTCD	AF TYTOC OCM	DNNFVSREDC	KRACAK A
	8) - KAVCSQEAMT	GPCRAVMPRT	TFDLSKGKCV	RF ITGCC COCM	RNNF BSEDYC	MAVCKAM
	9) RPDFCLEPPYT	<b>GPCKARIIR</b> Y	FYNAKAGLCQ	TF VY OCC RAK	RNNF KSAEDC	MRTCGGA
	10)CQLGYSA	<b>GPCMCMTSRY</b>	FYNGTSMACE	TF QY GGC MOM	GNNF VTEKEC	LQTC
	11) VAACNLPIVR	<b>GPCRAFIQLW</b>	AFDAVKGKCV	LF PYGGC QCM	CNKF YSEKEC	REYCGVP
15	12) - EVCCSEQAET	<b>GPCRAMISRW</b>	YFDVTEGKCA	PF TYGGC GCT	RNNF DTEEYC	MAVCGSA
	13)CKLPKDE	GTCROFILAW	YYDPNTKSCA	RFWYGGC GCM	enkf asokec	EKVC
	14) - PNVCAFPMER	GPCQTYMTRW	FFNFETGECE	LF AYGGC GCM	SNNF LRKEKC	exfcxf <b>t</b>

Where sequence number 1) is Bikunin (7-64) (SEQ ID NO: 4); sequence 2) is Bikunin (102-159) (SEQ ID NO: 6); sequence 3) is Tissue factor pathway inhibitor precursor 1 (SEQ ID NO: 18); sequence 4) is Tissue factor pathway inhibitor precursor 1 (SEQ ID NO: 19); sequence 5) is Tissue factor pathway inhibitor precursor (SEQ ID NO: 20); sequence 6) is Tissue factor pathway inhibitor precursor 2 (SEQ ID NO: 21); sequence 7) is Tissue factor pathway inhibitor precursor 2 (SEQ ID NO: 22); sequence 8) is Amyloid precursor protein homologue (SEQ ID NO: 23); sequence 9) is Aprotinin (SEQ ID NO: 24); sequence 10) is Inter-α-trypsin inhibitor precursor (SEQ ID NOs: 25); sequence 11) is Inter-α-trypsin inhibitor precursor (SEQ ID NOs: 26); sequence 12) is Amyloid precursor protein (SEQ ID NO: 27); sequence 13) is Collagen α-3(VI) precursor (SEQ ID NO: 28); and squence 14) is HKI-B9 (SEQ ID NO: 29).

It can be seen that Placental Bikunin (7-64) and (102-159) each have the same number (six) and spacing of cysteine residues as is found in members of the Kunitz class of serine protease inhibitors. The precise bonding of cysteine residues to form the three intrachain disulfide bonds is known and invarient for all previously known Kunitz family members (Laskowski, M et al., 1980, Ann. Rev. Biochem. 49:593-626). Based on this known bonding pattern and the fact that the folding of Placental Bikunin (7-64) and (102-159) into active protease inhibitors is accompanied by a mass reduction consistent with the formation of

three intrachain disulfide bonds (Examples 2 and 1), it is highly probable that the disulfide bonding within the Kunitz domains of Placental Bikunin occur between cysteine residues: C11 and C61; C20 and C44; C36 and C57; C106 and C156; C115 and C139; C131 and C152. Furthermore, this pattern of disulfide bonding is highly probable in larger forms of Placental Bikunin containing both Kunitz domains since such forms of the protein are also active serine protease inhibitors and because N-terminal sequencing (Example 7) of native Placental Bikunin for 50 cycles yielded a sequence that was silent at positions where the cysteine residues were expected.

The placental bikunin, isolated domains or other variants of the present invention may be produced by standard solid phase peptide synthesis using either t-Boc chemistry as described by Merrifield R.B. and Barany G., in: The peptides, Analysis, Synthesis, Biology, 2, Gross E. et al., Eds. Academic Press (1980) Chapter 1; or using F-moc chemistry as described by Carpino L.A., and Han G.Y., (1970) J. Amer Chem Soc., 92, 5748-5749, and illustrated in Example 2. Alternatively, expression of a DNA encoding the placental bikunin variant may be used to produce recombinant placental bikunin variants.

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The invention also relates to DNA constructs that encode the Placental bikunin protein variants of the present invention. These constructs may be prepared by synthetic methods such as those described in Beaucage S.L. and Caruthers M.H., (1981) Tetrahedron Lett, 22, pp1859-1862; Matteucci M.D and Caruthers M.H., (1981), J. Am. Chem. Soc. 103, p 3185; or from genomic or cDNA which may have been obtained by screening genomic or cDNA libraries with cDNA probes designed to hybridize with placental bikunin encoding DNA sequence. Genomic or cDNA sequence can be modified at one or more sites to obtain cDNA encoding any of the amino acid substitutions or deletions described in this disclosure.

The instant invention also relates to expression vectors containing the DNA constructs encoding the placental bikunin, isolated domains or other variants of the present invention that can be used for the production of recombinant placental bikunin variants. The cDNA should be connected to a suitable promoter sequence which shows transcriptional activity in the host cell moice, possess a suitable terminator and a poly-adenyiation signa.

CDNA encoding the piacental bikunin variant can be rused to a posignal peptide that will result in the protein encoded by the cDNA to undergo secretion. The signal peptide can be one that is recognized by the host organism. In the case of a mammalian host cell, the signal peptide can also be the natural signal peptide.

present in full length placental bikunin. The procedures used to prepare such vectors for expression of placental bikunin variants are well known in the art and are for example described in Sambrook et al., Molecular Cloning: A laboratory Manual, Cold Spring Harbor, New York, (1989).

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The instant invention also relates to transformed cells containing the DNA constructs encoding the placental bikunin, isolated domains or other variants of the present invention that can be used for the production of recombinant placental bikunin variants. A variety of combinations of expression vector and host organism exist which can be used for the production of the placental bikunin variants. Suitable host cells include baculovirus infected Sf9 insect cells, mammalian cells such as BHK, CHO, Hela and C-127, bacteria such as E. coli, and yeasts such as Saccharomyces cervisiae. Methods for the use of mammalian, insect and microbial expressions systems needed to achieve expression of placental bikunin are well known in the art and are described, for example, in Ausubel F.M et al., Current Protocols in Molecular Biology, John Wiley & Sons (1995), Chapter 16. For fragments of placental bikunin containing a single Kunitz inhibitor domain such as bikunin (7-64) and (102-159), yeast and E. coli expression systems are preferable, with yeast systems being most preferred. Typically, yeast expression would be carried out as described in US patent 5,164,482 for aprotinin variants and adapted in Example 5 of the present specification for placental bikunin (102-159). E.coli expression could be carried out using the methods described in US patent 5,032,573. Use of mammalian and yeast systems are most preferred for the expression of larger placental bikunin variants containing both inhibitor domains such as the variant bikunin (7-159).

DNA encoding variants of placental bikunin that possess amino acid substitution of the natural amino sequence can be prepared for expression of recombinant protein using the methods of Kunkel T.A., (1985) Proc. Natl. Acad. Sci USA 82: 488-492. Briefly, the DNA to be mutagenized is cloned into a single stranded bacteriophage vector such as M13. An oligonucleotide spanning the region to be changed and encoding the substitution is hybridized to the single stranded DNA and made double stranded by standard molecular biology techniques. This DNA is then transformed into an appropriate bacterial host and verified by dideoxynucleotide sequencing. The correct DNA is then cloned into the expression plasmid. Alternatively, the target DNA may be mutagenized by standard PCR techniques, sequenced, and inserted into the appropriate expression plasmid.

The following particular examples are offered by way of illustration, and not limitation, of certain aspects and preferred embodiments of the instant invention.

# 5 Example 1

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Preparation of synthetic placental bikunin (102-159)

Materials and methods/Reagents used. The fluorogenic substrate Tos-Gly-Pro-Lys-AMC was purchased from Bachem BioScience Inc (King of Prussia, PA). PNGB, Pro-Phe-Arg-AMC, Ala-Ala-Pro-Met-AMC, bovine trypsin (type III), human plasma kallikrein, and human plasmin were from Sigma (St. Louis, MO).

Recombinant aprotinin (Trasylol®) was from Bayer AG (Wuppertal, Germany). Pre-loaded Gln Wang resin was from Novabiochem (La Jolla, CA). Thioanisole, ethanedithiol and t-butyl methyl ether was from Aldrich (Milwaukee, WI).

# Quantification of functional placental bikunin (7-64) and (102-159)

The amount of trypsin inhibitory activity present in the refolded sample at various stages of purification was measured using GPK-AMC as a substrate. Bovine trypsin (200 pmoles) was incubated for 5 min at 37%C with bikunin (7-20 64) or (102-159), from various stages of purification, in buffer A (50 mM Hepes, pH 7.5, 0.1 M NaCl, 2 mM CaCl<sub>2</sub> and 0.01% triton X-100). GPK-AMC was added (20 µM final) and the amount of coumarin produced was determined by measuring the fluorescence (ex = 370 nm, em = 432 nm) on a Perkin-Elmer LS-50B fluorimeter over a 2 min. period. For samples being tested the % inhibition 25 for each was calculated according to equation 1; where  $R_0$  is the rate of fluorescence increase in the presence of inhibitor and  $R_1$  is the rate determined in the absence of added sample. One unit of activity for the inhibitor is defined as the amount needed to achieve 50% inhibition in the assay using the 30 conditions as described.

% inhibition =  $100 \times [1 - R_0/R_1]$  (1)

Biosystems model 420A peptide synthesizer using NMP-HBTU Fmoc chemistry. The peptide was synthesized on pre loaded Gln resin with an 8-fold excess of amino acid for each coupling. Cleavage and deprotection was

performed in 84.6% trifluoroacetic acid (TFA), 4.4% thioanisole, 2.2% ethanedithiol, 4.4% liquified phenol, and 4.4% H<sub>2</sub>O for 2 hours at room temperature. The crude peptide was precipitated, centrifuged and washed twice in t-butyl methyl ether. The peptide was purified on a Dynamax 60A C18 reverse-phase HPLC column using a TFA/acetonitrile gradient. The final preparation (61.0 mg) yielded the correct amino acid composition and molecular mass by Electrospray mass spectroscopy (MH+ =6836.1; calcd = 6835.5) for the predicted sequence:

YEEYCTANAV TGPCRASFPR WYFDVERNSC NNF1YGGCRG NKNSYRSEEA

CMLRCFRQ (SEQ ID NO: 6)

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Purification. Refolding of placental bikunin (102-159) was performed according to the method of Tam et al., (J. Am. Chem. Soc. 1991, 113: 6657-62). A portion of the purified peptide (15.2 mg) was dissolved in 4.0 ml of 0.1 M Tris, pH 6.0, and 8 M urea. Oxidation of the disulfides was accomplished by 15 dropwise addition of a solution containing 23% DMSO, and 0.1 M Tris, pH 6.0 to obtain a final concentration of 0.5 mg/ml peptide in 20% DMSO, 0.1 M Tris, pH 6.0, and 1 M urea. The solution was allowed to stir for 24 hr at 25℃ after which it was diluted 1:10 in buffer containing 50 mM Tris, pH 8.0, and 0.1 M NaCl. The material was purified using a kallikrein affinity column made by 20 covalently attaching 30 mg of bovine pancreatic kallikrein (Bayer AG) to 3.5 mls of CNBr activated Sepharose (Pharmacia) according to the manufacturers instructions. The refolded material was loaded onto the affinity column at a flow rate of 1 ml/min and washed with 50 mM Tris, pH 8.0, and 0.1 M NaCl until absorbance at 280 nm of the wash could no longer be detected. The 25 column was eluted with 3 volumes each of 0.2 M acetic acid, pH 4.0 and 1.7. Active fractions were pooled (see below) and the pH of the solution adjusted to 2.5. The material was directly applied to a Vydac C18 reverse-phase column (5 micron, 0.46 x 25 cm) which had been equilibrated in 22.5% acetonitrile in 0.1% TFA. Separation was achieved using a linear gradient of 22.5 to 40% acetonitrile 30 in 0.1% TFA at 1.0 ml/min over 40 min. Active fractions were pooled, lyophilized, redissolved in 0.1% TFA, and stored at -20°C until needed.

Results. Synthetic placental bikunin (102-159) was refolded using 20% DMSO as the oxidizing agent as described above, and purified by a 2-step purification protocol as shown below, to yield an active trypsin inhibitor (Table 1 below).

Table 1
Purification table for the isolation of synthetic placental bikunin (102-159)

Purification Step	Vol (ml)	mg/ml	mg	Units <sup>c</sup> (U)	SpA (U/mg)	Yield
8.0 M Urea	4.0	3.75 a	15.0	0	0	<del></del> -
20% DMSO	32.0	0.47 2	15.0	16,162	1,078	100
Kallikrein affinity	9.8	0.009 b	0.09	15,700	170,000	97
C18	3.0	0.013 ab	0.04	11,964	300,000	74

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<sup>a</sup>Protein determined by AAA.

<sup>b</sup>Protein determined by OD280 nm using the extinction coefficient determined for the purified protein (1.7 x 10<sup>4</sup> Lmol<sup>-1</sup> cm<sup>-1</sup>).

COne Unit is defined as the amount of material required to inhibit 50% of trypsin activity in a standard assay.

Chromatography of the crude refolded material over an immobilized bovine pancreatic kallikrein column selectively isolated 6.0% of the protein and 97% of the trypsin inhibitory activity present. Subsequent chromatography using C18 reverse-phase yielded a further purification of 2-fold, with an overall recovery of 74%. On RPHPLC, the reduced and refolded placental bikunin (102-159), exhibited elution times of 26.3 and 20.1 minutes, respectively. Mass spectroscopy analysis of the purified material revealed a molecular mass of 6829.8; a loss of 6 mass units from the starting material. This demonstrates the complete formation of the 3 disulfides predicted from the peptide sequence.

The isoelectric points of the purified, refolded synthetic placental bikunin (102-159) was determined using a Multiphor II Electrophoresis System (Pharmacia) run according to the manufacturers suggestions, together with pI standards, using a precast Ampholine® PAGplate (pH 3.5 to 9.5) and focused for 1.5 hrs. After staining, the migration distance from the cathodic edge of the gel to the different protein bands was measured. The pI of each unknown was determined by using a standard curve generated by a plot of the migration

value predicted from the amino acid sequence. This is lower than the value or 10.5 established for the pI of aprotinin. (Tenstad et al., 1994, Acta Physiol. Scand. 152: 33-50).

## Example 2

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# Preparation of synthetic placental bikunin (7-64)

Placental bikunin (7-64) was synthesized, refolded and purified essentially as described for placental bikunin (102-159) but with the following modifications: during refolding, the synthetic peptide was stirred for 30 hr as a solution in 20% DMSO at 25°C; purification by C18 RP-HPLC was achieved with a linear gradient of 25 to 45% acetonitrile in 0.1% TFA over 40 min (1ml/min). Active fractions from the first C18 run were reapplied to the column and fractionated with a linear gradient (60 min, 1 ml/min) of 20 to 40% acetonitrile in 0.1% TFA.

**Results.** The final purified reduced peptide exhibited an MH+=6563, consistent with the sequence:

15 IHDFCLVSKV VGRCRASMPR WWYNVTDGSC QLFVYGGCDG NSNNYLTKEE CLKKCATV (SEQ ID NO: 4)

The refolding and purification yielded a functional Kunitz domain that was active as an inhibitor of trypsin (Table 2 below).

Table 2A

Purification table for the isolation of synthetic placental bikunin (7-64)

Purification Step	Vol (ml)	mg/ml	mg	Units (U)	SpA (U/mg)	Yiek
8.0 M Urea	8.0	25	20.0	0	0	•
20% DMSO	64.0	0.31	20.0	68,699	3,435	100
Kall affirity pH 4.0	11.7	0.10	1.16	43,333	36,110	62
Kall affireity pH 1.7	9.0	0.64	5.8	4972	857	7.2
C18-1	4.6	0.14	0.06	21,905	350,143	31.9
C18-2	1.0	0.08	0.02	7,937	466,882	115

The purified refolded protein exhibited an MH+ = 6558, i.e.  $5\pm1$  mass units less than for the reduced peptide. This demonstrates that refolding caused the formation of at least one appropriate disulfide bond.

The pI of placental bikunin (7-64) was determined using the methods employed to determine the pI of placental bikunin (102-159). Placental bikunin (7-64) exhibited a pI that was much higher than the predicted value (pI = 7.9).

Refolded placental bikunin (7-64) migrated to the cathodic edge of the gel (pH 9.5) and an accurate pI could not be determined under these conditions.

# Continued Preparation of synthetic placental bikunin (7-64)

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Because the synthetic placental bikunin (7-64) may not have undergone complete deprotection prior to purification and refolding, refolding was repeated using protein which was certain to be completely deprotected. Placental bikunin (7-64) was synthesized, refolded and purified essentially as described for placental bikunin (102-159) but with the following modifications: during refolding, the synthetic peptide (0.27 mg/ml) was stirred for 30 hr as a solution in 20% DMSO at 25 C; purification by C18 RP-HPLC was achieved with a linear gradient of 22.5 to 50% acetonitrile in 0.1% TFA over 40 min (1 ml/min).

**Results.** The final purified reduced peptide exhibited an MH+ = 6567.5, consistent with the sequence:

IHDFCLVSKV VGRCRASMPRW WYNVTDGSC QLFVYGGCDG NSNNYLTKEE CLKKCATV (SEQ ID NO: 4)

The refolding and purification yielded a functional Kunitz domain that was as active as an inhibitor of trypsin (Table 2B below).

Table 2B
Purification table for the isolation of synthetic placental bikunin (7-64)

TABLE 2B						
Purification Step	Vol (ml)	mg/ml	mg	Units (U)	SpA (U/mg)	Yield
8.0 M Urea	4.9	2.1	10.5	0	0	•
20% DMSO	39.0	0.27	10.5	236,000	22,500	100
Kallikrein Affinity (pH 2)	14.5	0.3	0.43	120,000	279,070	50.9
C18 Reverse- Phase	0.2	1.2	0.24	70,676	294,483	30.0

The purified refolded protein exhibited an MH+ = 6561.2, i.e. 6.3 mass units less than for the reduced peptide. This demonstrates that refolding caused the armanon or the expected three distinge cond.

The pi or retolded placental bikunin (v-64) was determined using the inclined employed to determine the pI of placental bikunin (102-159). Refolded placental bikunin (7-64) exhibited a pI of 8.85, slightly higher than the predicted value (pI = 79).

## Example 3

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In vitro specificity of functional placental bikunin fragment (102-159)

Protenses. Bovine trypsin, human plasmin, and bovine pancreatic kallikrein quantitation was carried out by active site titration using p-nitrophenyl p'-guanidinobenzoate HCl as previously described (Chase,T., and Shaw, E., (1970) Methods Enzmol., 19: 20-27). Human kallikrein was quantitated by active site titration using bovine aprotinin as a standard and PFR-AMC as a substrate assuming a 1:1 complex formation. The K<sub>m</sub> for GPK-

AMC with trypsin and plasmin under the conditions used for each enzyme was 29 μM and 726 μM, respectively; the K<sub>m</sub> for PFR-AMC with human plasma kallikrein and bovine pancreatic kallikrein was 457 μM and 81.5 μM, respectively; the K<sub>m</sub> for AAPR-AMC with elastase was 1600 μM. Human tissue kallikrein (Bayer, Germany) quantification was carried out by active site titration using p'nitrophenyl p'-guanidinobenzoate HCl as previously described (Chase, T., and Shaw, E., (1970) Methods Enzmol. 19: 20-27).

Inhibition Kinetics: The inhibition of trypsin by placental bikunin (102-159) or aprotinin was measured by the incubation of 50 pM trypsin with placental bikunin (102-159) (0-2 nM) or aprotinin (0-3 nM) in buffer A in a total 20 volume of 1.0 ml. After 5 min. at 37°C, 15 µl of 2 mM GPK-AMC was added and the change in fluorescence (as above) was monitored. The inhibition of human plasmin by placental bikunin (102-159) and aprotinin was determined with plasmin (50 pM) and placental bikunin (102-159) (0-10 nM) or aprotinin (0-4 nM) in buffer containing 50 mM Tris-HCl (pH 7.5), 0.1 M NaCl, and 0.02% 25 triton x-100. After 5 min. incubation at 37°C, 25 µl of 20 mM GPK-AMC was added and the change in fluorescence monitored. The inhibition of human plasma kallikrein by placental bikunin (102-159) or aprotinin was determined using kallikrein (2.5 nM) and placental bikunin (102-159) (0-3 nM) or aprotinin (0-45 nM) in 50 mM Tris-HCl (pH 8.0), 50 mM NaCl, and 0.02% triton x-100. 30 After 5 min. at 37°C 15 µl of 20 mM PFR-AMC was added and the change in fluorescence monitored. The inhibition of bovine pancreatic kallikrein by placental bikunin (102-159) and aprotinin was determined in a similar manner with kallikrein (92 pM), placental bikunin (102-159) (0-1.6 nM) and aprotinin (0-14 pM) and a final substrate concentration of 100 µM. The apparent inhibition 35 constant Ki\* was determined using the nonlinear regression data analysis program Enzfitter software (Biosoft, Cambridge, UK): The kinetic data from

each experiment were analyzed in terms of the equation for a tight binding inhibitor:

$$V_i/V_0 = 1 - (E_0 + I_0 + K_i^* - [(E_0 + I_0 + K_i^*)^2 - 4 E_0 I_0]^{1/2})/2E_0$$
 (2)

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where  $V_i/V_O$  is the fractional enzyme activity (inhibited vs. uninhibited rate), and  $E_O$  and  $I_O$  are the total concentrations of enzyme and inhibitor, respectively. Ki values were obtained by correcting for the effect of substrate according to the equation:

$$K_i = K_i^* / (1 + [S_0] / K_m)$$
 (3)

(Boudier, C., and Bieth, J. G., (1989) Biochim Biophys Acta., 995: 36-41)

For the inhibition of human neutrophil elastase by placental bikunin (102-159) and aprotinin, elastase (19 nM) was incubated with placental bikunin (102-159) (150 nM) or aprotinin (0-7.5  $\mu$ M) in buffer containing 0.1 M Tris-HCl (pH 8.0), and 0.05% triton X-100. After 5 min at 37%C, AAPM-AMC (500  $\mu$ M or 1000  $\mu$ M) was added and the fluorescence measured over a two-minute period. Ki values were determined from Dixon plots of the form 1/V versus [I] performed at two different substrate concentrations (Dixon et al., 1979).

The inhibition of human tissue kallikrein by aprotinin, placental bikunin fragment (7-64) or placental bikunin fragment (102-159) was measured by the incubation of 0.35 nM human tissue kallikrein with placental bikunin (7-64) (0-40 nM) or placental bikunin (102-159) (0-2.5 nM), or aprotinin (0-0.5 nM) in a 1 ml reaction volume containing 50 mM Tris-HCl buffer pH 9.0, 50 mM NaCl, and 0.1% triton x-100. After 5 min. at 37°C, 5 ul of 2 mM PFR-AMC was added achieving 10 uM final and the change in fluorescence monitored. The Km for PFR-AMC with human tissue kallikrein under the conditions employed was 5.7 uM. The inhibition of human factor Xa (American Diagnostica, Inc, Greenwich, CT) by synthetic placental bikunin (102-159), recombinant placental bikunin, and aprotinin was measured by the incubation of 0.87 nM human factor Xa with ocreasing amounts of inhibitor in buffer containing 20 mM Tris (pH 7.5), 0.1 M Fact and 0.1% BSA. After a min of the so of 20 mm lock-AMC digma was added and the change in fluorescence monitored. The inhibition or numai. urokinase (Sigma) by Kunitz inhibitors was measured by the incubation of urokinase (2.7 ng) with inhibitor in a total volume of 1 ml buffer containing 50 mM Tris-HCl (pH 8.0), 50 mM NaCl, and 0.1% Triton x-100. After 5 min. at

37°C, 35 ul of 20 mM GGR-AMC (Sigma) was added and the change in fluorescence monitored. The inhibition of Factor XIa (from Enzyme Research Labs, Southbend, IN) was measured by incubating FXIa (0.1 nM) with either 0 to 800 nM placental bikunin (7-64), 0 to 140 nM placental bikunin (102-159) or 0 to 40 uM aprotinin in buffer containing 50 mM Hepes pH 7.5, 100 mM NaCl, 2 mM CaCl2, 0.01% triton x-100, and 1% BSA in a total volume of 1 ml. After 5 min at 37 C, 10 ul of 40 mM Boc-Glu(OBzl)-Ala-Arg-AMC (Bachem Biosciences, King of Prussia, PA) was added and the change in fluorescence monitored.

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10 Results: A direct comparison of the inhibition profiles of placental bikunin (102-159) and aprotinin was made by measuring their inhibition constants with various proteases under identical conditions. The K<sub>i</sub> values are listed in Table 3 below.

Table 3
Ki values for the inhibition of various proteases by bikunin (102-159)

Protesse (concentration)	bikunin (102-159) Ki (nM)	Aprotinin Ki (nM)	Substrate (concentration)	Km (mM)
Trypsin (48.5 pM)	0.4	0.8	GPK-AMC (0.03 mM)	0.022
Chymotrypsin (5 nM)	0.24	0.86	AAPF-pNA (0.08 mM)	0.027
Bovine Pancreatic Kallikrein (92.0 pM)	0.4	0.02	PFR-AMC (0.1 mM)	0.08
Human Plasma Kallikrein (2.5 nM)	0.3	19.0	PFR-AMC (0.3 mM)	0.46
Human Plasmin (50 pM)	1.8	13	GPK-AMC (0.5 mM)	0.73
Human Neutrophil Elastase (19 nM)	323.0	8500.0	AAPM-AMC (1.0 µM)	1.6
Factor XIIa	>300.0	12,000.0	PFR-AMC (0.2 µM)	0.35
Human Tissue Kallikrein (0.35 nM)	0.13	0.004	PFR-AMC (10 µM)	0.0057
factor Xa (0.67 nM)	274	NJ. at 3 µM	LGR-AMC (0.6 mM)	N.D.
urokinase	11000	4500	CCR-AMC (0.7 mM)	N.D.
factor XIa (0.1 nM)	15	288	E(OBz)AR-AMC (0.4 mM)	0.46

Placental bikunin (102-159) and aprotinin inhibit bovine trypsin and human plasmin to a comparable extent under the conditions employed. Aprotinin inhibited elastase with a Ki of 8.5 µM. Placental bikunin (102-159) inhibited elastase with a Ki of 323nM. The Ki value for the placental bikunin (102-159) inhibition of bovine pancreatic kallikrein was 20-fold higher than that of aprotinin inhibition. In contrast, placental bikunin (102-159) is a more potent

inhibitor of human plasma kallikrein than aprotinin and binds with a 56-fold higher affinity.

Because placental bikunin (102-159) is greater than 50 times more potent than Trasylol® as an inhibitor of kallikrein, smaller amounts of human placental bikunin, or fragments thereof (i.e. placental bikunin (102-159)) are needed than Trasylol® in order to maintain the effective patient doses of inhibitor in KIU. This reduces the cost per dose of the drug and reduces the likelihood of adverse nephrotoxic effects upon re-exposure of the medicament to patients. Furthermore, the protein is human derived, and thus much less immunogenic in man than aprotinin which is derived from cows. This results in significant reductions in the risk of incurring adverse immunologic events upon re-exposure of the medicament to patients.

## Example 4

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# 15 In vitro specificity of functional placental bikunin fragment (7-64)

In vitro specificity of functional human placental bikunin (7-64) was determined using the materials and methods as described in the Examples above.

20 Results: The table below shows the efficacy of placental bikunin (7-64) as an inhibitor of various serine proteases in vitro. Data is shown compared against data obtained for screening inhibition using either placental bikunin (102-159), or aprotinin (Trasylol®).

Table 4 A
Ki values for the inhibition of various proteases by bikunin(7-64)

TABLE 4A			
Protease (concentration)	bikunin(7-64) Ki (nM)	Aprotinin Ki (nM)	bikunin (102-159) Ki (nM)
Trypsin (48.5 pM)	0.17	0.8	0.4
Bovine Pancreatic Kallikrein (92.0 pM)	0.4	0.02	0.4
Human Plasma Kallikr / n (2.5 nM)	2.4	19.0	0.3
Human Plasmin	7 •	13	18
sovine chymotrypsin 5 nM)	Še		Production of the state of the
Factor XIIa	>300	12000	>300
elastase	>100	8500	323

The results show that the amino acid sequence encoding placental bikunin (7-64) can be refolded to obtain an active serine protease inhibitor that is effective against at least four trypsin-like serine proteases.

Table 4B below also shows the efficacy of refolded placental bikunin (7-64) as an inhibitor of various serine proteases *in vitro*. Refolded placental bikunin (7-64) was prepared from protein that was certain to be completely deprotected prior to purification and refolding. Data is shown compared against data obtained for screening inhibition using either placental bikunin (102-159), or aprotinin (Trasylol®).

Table 4B Ki values for the inhibition of various proteases by refolded bikunin (7-54)

Protease (concentration)	bikunin (7-64) Ki (nM)	Aprotinin Ki (nM)	bikunin (102-159) Ki (nM)
Trypsin (50 pM)	0.2	0.8	0.3
Human Plasma Kallikrein (0.2 nM)	0.7	19.0	0.7
Human Plasmin (50 pM)	3.7	1.3	1.8
Factor XIIa	not done	12,000	4,500
Factor XIa (0.1 nM)	200	288	15
Human Tissue Kallikrein	2.3	0.004	0.13

Suprisingly, placental bikunin (7-64) was more potent than aprotinin at inhibiting human plasma kallikrein, and at least similar in efficacy as a plasmin inhibitor. These data show that placental bikunin (7-64) is at least as effective as aprotinin, using *in vitro* assays, and that one would expect better or similar potency *in vivo*.

# Example 5

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# Expression of placental bikunin variant (102-159) in yeast

The DNA sequence encoding placental bikunin 102-159 (SEQ ID NO: 6) was generated using synthetic oligonucleotides. The final DNA product consisted (5' to 3') of 15 nucleotides from the yeast α-mating factor propeptide sequence fused to the in-frame cDNA sequence encoding placental bikunin (102-159), followed by an in-frame stop codon. Upon cloning into a yeast expression vector pS604, the cDNA would direct the expression of a fusion protein comprising an N-terminal yeast α-mating factor propeptide fused to the 58 amino acid sequence of placental bikunin (102-159). Processing of this fusion

protein at a KEX-2 cleavage site at the junction between the  $\alpha$ -mating factor and Kunitz domain was designed to liberate the Kunitz domain at its native N-terminus.

A 5' sense oligonucleotide of the following sequence and containing a HindIII site for cloning was synthesized:

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GAA GGG GTA AGC TTG GAT AAA AGA TAT GAA GAA TAC TGC ACC GCC AAC GCA GTC GTC ACC GGC GCA GTC GTG GAG AGG (SEQ ID NO: 42)

A 3' antisense oligonucleotide of the following sequence and containing both a BamHI site for cloning and a stop codon was synthesized:

CGC GGA TCC CTA CTG GCG GAA GCA GCG GAG CAT GCA GGC CTC CTC
AGA GCG GTA GCT GTT CTT ATT GCC CCG GCA GCC TCC ATA GAT GAA
GTT ATT GCA GGA GTT CCT CTC CAC GTC AAA GTA CCA GCG
(SEQ ID NO: 43)

The oligonucleotides were dissolved in 10 mM Tris buffer pH 8.0

containing 1 mM EDTA, and 12 ug of each oligo were added combined and brought to 0.25M NaCl. To hybridize, the oligonucleotides were denatured by boiling for 5 minutes and allowed to cool from 65°C to room temp over 2 hrs. Overlaps were extended using the Klenow fragment and digested with HindIII and BamHI. The resulting digested double stranded fragment was cloned into pUC19 and sequence confirmed. A clone containing the fragment of the correct sequence was digested with BamHI/HindIII to liberate the bikunin containing fragment with the following + strand sequence:

GAA GGG GTA AGC TTG GAT AAA AGA TAT GAA GAA TAC TGC ACC GCC

AAC GCA GTC ACT GGG CCT TGC CGT GCA TCC TTC CCA CGC TGG TAC

TTT GAC GTG GAG AGG AAC TCC TGC AAT AAC TTC ATC TAT GGA GGC

TGC CGG GGC AAT AAG AAC AGC TAC CGC TCT GAG GAG GCC TGC ATG

CTC CGC TGC TTC CGC CAG TAG GGA TCC (SEQ ID.: 44)

which was then gel purified and ligated into BamHI/HindIII cut pS604. The Lation maxture was Extracted to offerior adortorors and curred strains of column. The ligation product was directed transformed into read strains SC101 and WHL341 and plated on ura selection plates. Twelve colorues from each strain were re-streaked on ura drop out plates. A single colony was inoculated into 2 ml of ura DO media and grown over night at 30°C. Cells were

pelleted for 2 minutes at 14000x g and the supernatants evaluated for their content of placental bikunin (102-159).

Detection of expression of placental bikunin (102-159) in transformed yeast

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Firstly, the supernatants (50 ul per assay) were evaluated for their capacity to inhibit the *in vitro* activity of trypsin using the assay methods as described in Example 1 (1 ml assay volume). An un-used media only sample as well as a yeast clone expressing an inactive variant of aprotinin served as negative controls. A yeast clone expressing natural aprotinin served as a positive control and is shown for comparison.

The second method to quantify placental bikunin (102-159) expression exploited use of polyclonal antibodies (pAbs) against the synthetic peptide to monitor the accumulation of the recombinant peptide using Western blots. These studies were performed only with recombinants derived from strain SC101, since these produced greater inhibitory activity than recombinants derived from strain WHL341.

To produce the pAb, two 6-8 week old New Zealand White female rabbits (Hazelton Research Labs, Denver, Pa) were immunized on day zero with 250 ug of purified reduced synthetic placental bikunin (102-159), in Complete Freund's adjuvant, followed by boosts on days 14, 35 and 56 and 77 each with 125 ug of the same antigen in Incomplete Freund's adjuvant. Antiserum used in the present studies was collected after the third boost by established procedures. Polyclonal antibodies were purified from the antiserum over protein A.

Colonies 2.4 and 2.5 from transformation of yeast SC101 (Figure 8) as well as an aprotinin control were grown overnight in 50 ml of ura DO media at 30°C. Cells were pelleted and the supernatant concentrated 100-fold using a Centriprep 3 (Amicon, Beverly, MA) concentrator. Samples of each (30 µl) were subjected to SDS-PAGE on 10-20% tricine buffered gels (Novex, San Diego, CA) using the manufacturers procedures. Duplicate gels were either developed with a silver stain kit (Integrated Separation Systems, Nantick, MA) or transferred to nitrocellulose and developed with the purified polyclonal antibody elicited to synthetic bikunin (102-159). Alkaline-phosphatase conjugated goat anti-rabbit antibody was used as the secondary antibody according to the manufacturer's directions (Kirkegaard and Perry, Gaithersburg, MD).

Fermentation broth from a 1L culture of SC101 strain 2.4 was harvested by centrifugation (4,000 g x 30 min.) then applied to a 1.0 ml column of anhydrochymotrypsin-sepharose (Takara Biochemical Inc., CA), that was previously equilibrated with 50 mM Hepes buffer pH 7.5 containing 0.1M NaCl, 2 mM CaCl<sub>2</sub> and 0.01% (v/v) triton X-100. The column was washed with the same buffer but containing 1.0 M NaCl until the A280nm declined to zero, whereupon the column was eluted with 0.1M formic acid pH 2.5. Eluted fractions were pooled and applied to a C18 column (Vydac, 5um, 4.6 x 250 mm) previously equilibrated with 0.1% TFA, and eluted with a 50 min. linear gradient of 20 to 80% acetonitrile in 0.1% TFA. Fractions containing placental bikunin (102-159) were pooled and re-chromatographed on C18 employing elution with a linear 22.5 to 50% acetonitrile gradient in 0.1% TFA.

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Results. Figure 8 shows the percent trypsin activity inhibited by twelve colonies derived from the transformation of each of strains SC101 and WHL341. The results show that all twelve colonies of yeast strain SC101 transformed with the trypsin inhibitor placental bikunin (102-159) had the ability to produce a substantial amount of trypsin inhibitory activity compared to the negative controls both of which showed no ability to inhibit trypsin. The activity is therefore related to the expression of a specific inhibitor in the placental bikunin variant (102-159) transformed cells. The yeast WHL341 samples contained minimal trypsin inhibitory activity. This may be correlated to the slow growth observed with this strain under the conditions employed.

Figure 9 shows the SDS-PAGE and western analysis of the yeast SC101 supernatants. Silver stained SDS-PAGE of supernatants derived from recombinant yeasts 2.4 and 2.5 expressing placental bikunin (102-159) as well as from the yeast expressing aprotinin yielded a protein band running at approximated 6 kDa, corresponding to the size expected for each recombinant Kunitz inhibitor domain. Western analysis showed that the 6 kDa bands expressed by stains 2.4. and 2.5 reacted with the pAb elicited to placental bikunin (102-159). The same 6 kDa band in the aprotinin control did not react of the same antibody demonstrating the specificity of the antibody demonstrating the specificity of the antibody.

The final preparation of placental bikunin C-terminal domain was highly pure by silver-stained SDS-PAGE (Figure 10). The overall recovery of broth-derived trypsin inhibitory activity in the final preparation was 31%. N-terminal

sequencing of the purified inhibitor indicated that 40% of the protein is correctly processed to yield the correct N-terminus for placental bikunin (102-159) while about 60 % of the material contained a portion of the yeast α-mating factor. The purified material comprised an active serine protease inhibitor exhibiting an apparent Ki of 0.35 nM for the *in vitro* inhibition of plasma kallikrein.

In conclusion, the accumulation both of a protease inhibitor activity and a protein immunochemically related to synthetic bikunin (102-159) in fermentation broth as well as the isolation of placental bikunin (102-159) from one of the transformed lines provided proof of expression of placental bikunin in the recombinant yeast strains described herein, showing for the first time the utility of yeasts for the production of placental bikunin fragments.

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Additional constructs were prepared in an effort to augment the expression level of the Kunitz domain contained within placental bikunin 102-159, as well as to increase the yield of protein with the correct N-terminus. We hypothesized that the N-terminal residues of placental bikunin 102-159 (YEEY—) may have presented a cleavage site that is only poorly recognized by the yeast KEX-2 protease that enzymically removes the yeast a-factor pro-region. Therefore, we prepared yeast expression constructs for the production of placental bikunin 103-159 (N-terminus of EEY...), 101-159 (N-terminus of NYEEY...) and 98-159 (DMFNYEEY...) in order to modify the P' subsites surrounding the KEX-2 cleavage site. To attempt to augment the levels of recombinant protein expression, we also used the yeast preferred codons rather than mammalian preferred codons in preparing some of the constructs described below. The constructs were essentially prepared as described above for placental bikunin 102-159 (defined as construct #1) but with the following modifications:

Construct #2 placental bikunin 103-159, yeast codon usage A 5' sense oligonucleotide

and 3' antisense oligonucleotide

ACTGGATCCT CATTGGCGAA AACATCTCAA CATACAGGCT TCTTCAGATC TGTAAGAATT TTTATTACCT CTACAACCAC CGTAAATAAA ATTATTACAA GAATTTCTTT CAACATCAAA GTACCATCT (SEQ ID NO: 56)

- were manipulated as described for the production of an expression construct (construct #1 above) for the expression of placental bikunin 102-159
  - Construct #3 placental bikunin 101-159, yeast codon usage A 5' sense oligonucleotide

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GAAGGGGTAA GCTTGGATAA AAGAAATTAC GAAGAATACT GTACTGCTAA TGCTGTTACT GGTCCATGTA GAGCTTCTTT TCCAAGATGG TACTTTGATG TTGAAAGA (SEQ ID NO: 57)

and the same 3' antisense oligonucleotide as used for construct #2, were manipulated as described for the production of an expression construct (construct #1 above) for the expression of placental bikunin 102-159.

Construct #4 placental bikunin 98-159, yeast codon usage A 5' sense oligonucleotide

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GAAGGGGTAA GCTTGGATAA AAGAGATATG TTTAATTACG AAGAATACTG TACTGCTAAT GCTGTTACTG GTCCATGTAG AGCTTCTTTT CCAAGATGGT ACTTTGATGT TGAAAGA (SEQ ID NO: 58)

and the same 3' antisense oligonucleotide as used for construct #2, were manipulated as described for the production of an expression construct (construct #1 above).

Yeast strain SC101 (MATa, ura 3-52, suc 2) was transformed with the plasmids containing each of the above cDNAs, and proteins were expressed using the methods that were described above for the production of placental bikunin 102-159 with human codon usage. Approximately 250 ml of each yeast culture was

above. The relative amount of trypsin inhibitory activity in the applysate, the amount of purified protein recovered and the N-terminal sequence of the purified protein were determined and are listed below in Table 7.

Table 7
Relative production levels of different proteins containing the C-terminal Kunitz domain of placental bikunin

Cons	truct	Relative conc. of inhibitor in applysate	N-terminal amount (pmol)	sequence sequence	Comments
#2	103-159	none detected	none	none	no expression
#3	101-159	25 % inhibition	none	none	low expression
#4	98-15 <del>9</del>	93 % inhibition	910	DMFNYE-	good expression correct product
#1	102-159	82 % inhibition	480	AKEEGV-	expression of active incorrectly processed

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The results show that placental bikunin fragments of different lengths that contain the C-terminal Kunitz domain show wide variation in capacity to express functional secreted protein. Constructs expressing fragments 101-159 and 103-159 yielded little or low enzymic activity in the supernatants prior to purification, and N-terminal sequencing of 0.05 ml aliquots of each purified fraction yielded undetectable amounts of inhibitor. On the other hand expression either of placental bikunin 102-159 or 98-159 yielded significant amounts of protease activity prior to purification. N-terminal sequencing however showed that the purified protein recovered from expression of 102-159 was once again largely incorrectly processed, exhibiting an N-terminus consistent with processing of the majority of the pre-protein at a site within the yeast  $\alpha$ -mating factor pro-sequence. The purified protein recovered from expression of placental bikunin 98-159 however was processed entirely at the correct site to yield the correct N-terminus. Furthermore, nearly twice as much protein was recovered as compared to the recovery of placental bikunin 102-159. Placental bikunin 98-159 thus represents a preferred fragment length for the production of the C-terminal Kunitz domain of placental bikunin by the  $\alpha$ mating factor pre-pro sequence / KEX-2 processing system of S. cerevisiae,

## Example 6

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# Alternative procedure for yeast expression

The 58 amino acid peptide derived from the R74593 translation product can also be PCR amplified from either the R87894-R74593 PCR product cloned into the TA vector<sup>TM</sup> (Invitrogen, San Diego, CA) after DNA sequencing or from human placental cDNA. The amplified DNA product will consist of 19 nucleotides from the yeast  $\alpha$ -mating factor leader sequence mated to the R74593 sequence which codes for the YEEY--CFRQ (58 residues) so as to make the translation product in frame, constructing an  $\alpha$ -mating factor/Kunitz domain fusion protein. The protein sequence also contains a kex 2 cleavage which will liberate the Kunitz domain at its native N-terminus.

The 5' sense oligonucleotide which contains a HindIII site for cloning will contain the following sequence:

15 GCCAAGCTTG GATAAAAGAT ATGAAGAAT ACTGCACCGC CAACGCA (SEQ ID NO: 30)

The 3' antisense oligonucleotide contains a BamHI site for cloning as well as a stop codon and is of the following sequence:

20 GGGGATCCTC ACTGCTGGCG GAAGCAGCGG AGCAT (SEQ ID NO: 31)

The full 206 nucleotide cDNA sequence to be cloned into the yeast expression vector is of the following sequence:

25 CCAAGCTTGG ATAAAAGATA TGAAGAATAC TGCACCGCCA ACGCAGTCAC TGGGCCTTGC CGTGCATCCT TCCCACGCTG GTACTTTGAC GTGGAGAGGA ACTCCTGCAA TAACTTCATC TATGGAGGCT GCCGGGGCAA TAAGAACAGC TACCGCTCTG AGGAGGCCTG CATGCTCCGC TGCTTCCGCC AGCAGTGAGG ATCCCC (SEQ ID NO: 32)

After PCR amplification, this DNA will be digested with HindIII, BamHI and cloned into the yeast expression vector pMT15 (see US patent 5,164,482.

bami-II. The resulting plasmid vector is used to transform yeast strain 50.00 using the methods described in US patent 5,164,482. The URA 3+ yeast transformants are isolated and cultivated under inducing conditions. The yield of recombinant Placental bikunin variants is determined according to the

amount of trypsin inhibitory activity that accumulated in the culture supernatants over time using the in vitro assay method described above. Fermentation broths are centrifuged at 9000 rpm for 30 minutes. The supernatant is then filtered through a 0.4 then a 0.2 µm filter, diluted to a conductivity of 7.5 ms, and adjusted to pH 3 with citric acid. The sample is then batch absorbed onto 200 ml of S-sepharose fast flow (Pharmacia) in 50 mM sodium citrate pH 3 and stirred for 60 min. The gel is subsequently washed sequentially with 2 L of each of: 50 mM sodium citrate pH 3.0; 50 mM Tris-HCL pH 9.0; 20 mM HEPES pH 6.0. The washed gel is transferred into a suitable column and eluted with a linear gradient of 0 to 1 M sodium chloride in 20 mM HEPES pH 6.0. Eluted fractions containing in vitro trypsin inhibitory activity are then pooled and further purified either by a) chromatography over a column of immobilized anhydrotrypsin (essentially as described in Example 2); b) by chromatography over a column of immobilized bovine kallikrein; or c) a combination of conventional chromatographic steps including gel filtration and/or anion-exchange chromatography.

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# Example 7 Isolation and characterization of native human placental bikunin from placenta

Bikunin protein was purified to apparent homogeniety from whole frozen placenta (Analytical Biological Services, Inc, Wilmington, DE). The placenta (740 gm) was thawed to room temperature and cut into 0.5 to 1.0 cm pieces, placed on ice and washed with 600 ml PBS buffer. The wash was decanted and 240 ml of placenta pieces placed into a Waring blender. After adding 300 ml of buffer consisting of 0.1 M Tris (pH 8.0), and 0.1 M NaCl, the mixture was blended on high speed for 2 min, decanted into 750.0 ml centrifuge tubes, and placed on ice. This procedure was repeated until all material was processed. The combined slurry was centrifuged at 4500 x g for 60 minutes at 4°C. The supernatant was filtered through cheese cloth and the placental bikunin purified using a kallikrein affinity column made by covalently attaching 70 mg of bovine pancreatic kallikrein (Bayer AG) to 5.0 mls of CNBr activated Sepharose (Pharmacia) according to manufacturers instruction. The material was loaded onto the affinity column at a flow rate of 2.0 ml/min and washed with 0.1 M Tris (pH 8.0), 0.1 M NaCl until absorbance at 280 nm of the wash could no longer be detected. The column was further washed with 0.1 M Tris (pH 8.0), 0.5 M NaCl and then eluted with 3 volumes of 0.2 M acetic acid, pH 4.0. Fractions containing kallikrein and trypsin inhibitory (see below)

activity were pooled, frozen, and lyophilized. Placental bikunin was further purified by gel-filtration chromatography using a Superdex 75 10/30 (Pharmacia) column attached to a Beckman System Gold HPLC system. Briefly, the column was equilibrated in 0.1 M Tris, 0.15 M NaCl, and 0.1% Triton X-100 at a flow rate of  $0.5 \, \text{ml/min}$ . The lyophilized sample was reconstituted in  $1.0 \, \text{ml}$ of 0.1 M Tris, pH 8.0 and injected onto the gel-filtration column in 200  $\mu$ l aliquots. Fractions were collected (0.5 ml) and assayed for trypsin and kallikrein inhibitory activity. Active fractions were pooled, and the pH of the solution adjusted to 2.5 by addition of TFA. The material was directly applied to a Vydac C18 reverse-phase column (5 micron, 0.46 x 25 cm) which had been equilibrated in 20% acetonitrile in 0.1 %TFA. Separation was achieved using a linear gradient of 20 to 80% acetonitrile in 0.1% TFA at 1.0 ml/min over 50 minutes after an initial 20 minute wash at 20% acetonitrile in 0.1% TFA. Fractions (1ml) were collected and assayed for trypsin and kallikrein inhibitory activity. Fractions containing inhibitory activity were concentrated using a speed-vac concentrator (Savant) and subjected to N-terminal sequence analysis.

# Functional assays for Placental Bikunin:

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Identification of functional placental bikunin was achieved by measuring its ability to inhibit bovine trypsin and human plasma kallikrein. Trypsin inhibitory activity was performed in assay buffer (50 mM Hepes, pH 7.5, 0.1 M NaCl, 2.0 mM CaCl2, 0.1% Triton x-100) at room temperature in a 96-well microtiter plate (Perkin Elmer) using Gly-Pro-Lys-Aminomethylcoumarin as a substrate. The amount of coumarin produced by trypsin was determined by measuring the fluorescence (ex = 370 nm, em = 432 nm) on a Perkin-Elmer LS-50B fluorimeter equipped with a plate reader. Trypsin (23 µg in 100 µl buffer) was mixed with 20 µl of the sample to be tested and incubated for 10 minutes at 25°C. The reaction was started by the addition of 50 µl of the substrate GPK-AMC (33 µM final) in assay buffer. The fluorescence intensity was measured and the % inhibition for each fraction was determined by:

" inhihition - 100 x [1 Fo/F1]

where Fo is the fluorescence of the unknown and F1 is the fluorescence of the trypsin only control. Kallikrein inhibitory activity of the fractions was similarly measured using 7.0 nM kallikrein in assay buffer (50 mM Tris, pH 8.0, 50 mM NaCl, 0.1% triton x-100) and 66.0 µM Pro-Phe-Arg-AMC as a substrate

## Determination of the in vitro specificity of placental bikunin

The *In vitro* specificity of native human placental bikunin was determined using the materials and methods as described in the preceding examples above. Placental bikunin was quantified by active site titration against a known concentration of trypsin using GPK-AMC as a substrate to monitor the fraction of unbound trypsin.

### Protein Sequencing

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The 1 ml fraction (C18-29 Delaria) was reduced to 300 ml in volume, on a Speed Vac, to reduce the amount of organic solvent. The sample was then loaded onto a Hewlett-Packard miniature biphasic reaction column, and washed with 1 ml of 2% trifluoroacetic acid. The sample was sequenced on a Hewlett-Packard Model G1005A protein sequencing system using Edman degradation. Version 3.0 sequencing methods and all reagents were supplied by Hewlett-Packard. Sequence was confirmed for 50 cycles.

**Results.** Placental Bikunin was purified to apparent homogeniety by sequential kallikrein affinity, gel-filtration, and reverse-phase chromatography (see purification table below):

Table 5
Purification table for native Placental Bikunin (1-179)

TABLE 5					
Step	Vol (ml)	OD 280 (/ml)	OD 280	Units <sup>a</sup> (U)	Units/OD 280
Placenta Supernatant	1800.0	41.7	75,060	3,000,000	40.0
Kallikrein Affinity pH 4.0	20.0	0.17	3.36	16,000	4,880
Kallikrein Affinity pH 1.7	10.2	0.45	4.56	12,000	2,630
Superdex 75	15.0	0.0085	0.13	3,191	24,546

25 One Unit is defined as that amount which inhibits 50% of trypsin activity in a standard assay.

The majority of the kallikrein and trypsin inhibitory activity eluted from the kallikrein affinity column in the pH 4.0 elution. Subsequent gel-filtration chromatography (Figure 5) yielded a peak of kallikrein and trypsin inhibitory activity with a molecular weight range of 10 to 40 kDa as judged by a standard

curve generated by running molecular weight standards under identical conditions. Reverse-phase C18 chromatography (Figure 6) yielded 4 peaks of inhibitory activity with the most potent eluting at approximately 30 % acetonitrile. The activity associated with the first peak to elute from C18 (fraction 29) exhibited an amino acid sequence starting with amino acid 1 of the predicted amino acid sequence of placental bikunin (ADRER...; SEQ ID NO: 1), and was identical to the predicted sequence for 50 cycles of sequencing (underlined amino acids in Figure 3). Cysteine residues within this sequence stretch were silent as expected for sequencing of oxidized protein. The cysteine residues at amino acid positions 11 and 20 of mature placental bikunin were later identified from sequencing of the S-pyridylethylated protein whereupon PTH-pyridylethyl-cysteine was recovered at cycles 11 and 20.

Interestingly, the asparagine at amino acid residue number 30 of the sequence (Figure 3) was silent showing that this site is likely to be glycosylated. Fraction 29 yielded one major sequence corresponding to that of placental bikunin starting at residue #1 (27 pmol at cycle 1) plus a minor sequence (2 pmol) also derived from placental bikunin starting at residue 6 (SIHD...). This shows that the final preparation sequenced in fraction 29 is highly pure, and most likely responsible for the protease inhibitory activity associated with this fraction (Figure 6).

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Accordingly, the final preparation of placental bikunin from C18 chromatography was highly pure based on a silver-stained SDS-PAGE analysis (Figure 7), where the protein migrated with an apparent Mr of 24 kDa on a 10 to 20 % acrylamide tricine gel (Novex, San Diego, CA) calibrated with the following molecular weight markers: insulin (2.9 kDa); bovine trypsin inhibitor (5.8 kDa); lysozyme (14.7 kDa); β-lactaglobulin (18.4 kDa); carbonic anhydrase (29 kDa); and ovalbumin (43 kDa). The above size of placental bikunin on SDS-PAGE is consistent with that predicted from the full length coding sequence (Figure 4F).

As expected based on the N-terminal sequencing results described above, the purified protein reacted with an antibody elicited to placental bikunin (7-64) to yield a banú with the same Mr (Figure 12A) as observed for the purified preparation

corresponding to the full length protein was not observed. Rather, a fragment that co-migrated with synthetic bikunin (102-159) of approximately 6 kDa was observed. The simplest interpretation of these results is that the purified preparation had

undergone degradation subsequent to purification to yield an N-terminal fragment comprising the N-terminal domain and a C-terminal fragment comprising the C-terminal domain. Assuming that the fragment reactive against antiserum to placental bikunin (7-64) is devoid of the C-terminal end of the full length protein, the size (24 kDa) would suggest a high state of glycosylation.

Table 6. below shows the potency of *in vitro* inhibition of various serine proteases by placental bikunin. Data are compared with that obtained with aprotinin (Trasylol®).

Table 6
Ki values for the inhibition of various proteases by placental bikunin

TABLE 6 Protease (concentration)	Placental Bikunin Ki (nM)	Aprotinin Ki (nM)
Trypsin (48.5 pM)	0.13	0.8
Human Plasmin (50 pM)	1.9	1.3

The results show that placental bikunin isolated from a natural source (human placenta) is a potent inhibitor of trypsin-like serine proteases.

# Example 8

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Expression pattern of placental bikunin amongst different human organs and tissues

A multiple tissue northern was purchased from Clontech which contained 2 µg of polyA+ RNA from human heart, brain, placenta, lung, liver, skeletal muscle, kidney, and pancreas. Two different cDNA probes were used:

1) a gel purified cDNA encoding placental bikunin (102-159); 2) the 780 base pair PCR-derived cDNA (Figure 4E) liberated from a TA clone by digestion with EcoRI and gel purified. Each probe was labeled using <sup>32</sup>P-dCTP and a random priming labeling kit from Boehringer Mannheim Biochemicals (Indiana), then used to hybridize to the multiple tissue northern according to the manufacturers specifications. Autoradiographs were generated using Biomax film with an 18 hr exposure time, and developed using a Umax Scanner and scanned using Adobe Photoshop.

Results. The pattern of tissue expression observed using a placental bikunin (102-159) probe (Figure 11A) or a larger probe containing both Kunitz

domains of placental bikunin (Figure 11B) was essentially the same as might be expected. The placental bikunin mRNA was most abundant in pancreas and placenta. Significant levels were also observed in lung, brain and kidney, while lower levels were observed in heart and liver, and the mRNA was undetectable in skeletal muscle. The transcript size was 1.95 kilobases in all cases, in close agreement with the predicted size of placental bikunin deduced both from EST overlay and cloning of full length cDNA described in preceding sections.

The broad tissue distribution of the mRNA shows that placental bikunin is broadly expressed. Since the protein also contains a leader sequence it would have ample exposure to the human immune system, requiring that it become recognized as a self protein. Additional evidence for a broad tissue distribution of placental bikunin mRNA expression was derived from the fact that some of the EST entries with homology to placental bikunin (Figure 4B) were derived from human adult and infant brain, and human retina, breast, ovary, olfactory epithelium, and placenta. It is concluded therefore that administration of the native human protein to human patients would be unlikely to elicit an immune response.

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Interestingly, the expression pattern of placental bikunin is somewhat reminiscent of that for bovine aprotinin which is found in high levels in bovine lung and pancreas. To further elucidate the expression pattern of placental bikunin, RT-PCR of total RNA from the following human cells was determined: un-stimulated human umbilical vein endothelial cells (HUVECs), HK-2 (line derived from kidney proximal tubule), TF-1 (erythroleukemia line) and phorbolester (PMA)-stimulated human peripheral blood leukocytes. The probes used:

CACCTGATCGCGAGACCCC (sense; SEQ ID NO: 59); CTGGCGGAAGCAGCGGAGCATGC (antisense; SEQ ID NO: 60),

were designed to amplify a 600 b.p placental bikunin encoding cDNA fragment. Comparisons were normalized by inclusion of actin primers to amplify an 800 b.p. actin fragment. Whereas the 800 b.p. fragment identified on agarose gels at netrodium promide was or equal intensity and ranes are notice. The control of the control of the other cell lines. We conclude that placental bikunin is not expressed in at least some endothelial cells but is expressed in some leukocyte populations.

Example 9

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Purification and properties of Placental Bikunin (1-170) highly purified from a Baculovirus / Sf9 expression system

A large fragment of Placental bikurin containing both Kunitz domains (Placental Bikurin 1-170) was expressed in Sf9 cells as follows. Placental bikurin cDNA obtained by PCR (Figure 4E) and contained within a TA vector (see previous Examples) was liberated by digestion with HindIII and Xba1 yielding a fragment flanked by a 5' XbaI site and 3' HindIII site. This fragment was gel purified and then cloned into the M13mp19 vector (New England Biolabs, Beverly, MA). In vitro mutagenesis (Kunkel T.A., (1985) Proc. Natl. Acad. Sci. USA, 82: 488-492) was used to generate a Pst1 site 3' to the XbaI site at the 5' end, but 5' to the sequence encoding the ATG start site, natural placental bikunin signal peptide and mature placental bikunin coding sequence. The oligonucleotide used for the mutagenesis had the sequence:

5' CGC GTC TCG GCT GAC CTG GCC CTG CAG ATG GCG CAC GTG TGC GGG 3' (SEQ ID NO: 61)

A stop codon (TAG) and Bgill / Xmal site was similarly engineered at the 3' end of the cDNA using the oligonucleotide:

5' CTG CCC CTT GGC TCA AAG TAG GAA GAT CTT CCC CCC GGG GGG GTG GTT CTG GCG GGG CTG 3' (SEQ ID NO: 62).

The stop codon was in frame with the sequence encoding placental bikunin and caused termination immediately following the Lysine at amino acid residue 170, thus encoding a truncated placental bikunin fragment devoid of the putative transmembrane domain. The product from digestion with Pst1 and BglII was isolated and cloned into the BacPac8 vector for expression of Placental bikunin fragment (1-170) which contains both Kunitz domains but which is truncated immediately N-terminal to the putative transmembrane segment.

The expression of Bikunin by Sf-9 insect cells was optimal at a multiplicity of infection of 1 to 1 when the medium was harvested at 72 h post infection. After harvesting, the baculovirus cell culture supernatant (2L) was adjusted to pH 8.0 by the addition of Tris-HCl. Bikunin was purified by chromatography using a 5 ml bovine pancreatic kallikrein affinity column as previously described in Example 7 for

the purification of native placental bikunin from placenta. Eluted material was adjusted to pH 2.5 with TFA and subjected to chromatography on a C18 reverse-phase column (1.0 x 25 cm) equilibrated in 10% acetonitrile in 0.1% TFA at a flow rate of 1 ml/min. The bikunin was eluted with a linear gradient of 10 to 80% acetonitrile in 0.1% TFA over 40 min. Active fractions were pooled, lyophilized, redissolved in 50 mM Hepes (pH 7.5), 0.1 M NaCl, 2 mM CaCl2, and 0.1% triton x-100, and stored at -20°C until needed. The concentration of recombinant bikunin was determined by amino acid analysis.

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10 Results. Recombinant bikunin was purified from baculovirus cell culture supernatant using a 2-step purification protocol as shown below, to yield an active trypsin inhibitor (Table 8 below).

Table 8
Purification of recombinant bikunin from transformed culture supernatant

TABLE 8						
Purification Step	Vol (ml)	OD 280/ml	OD 280 total	Units (U)	Specific activity (U/OD)	
Supernatant	2300.0	9.0	20,700	6,150,0 <b>00</b>	297	
Kallikrein affinity	23.0	0.12	2.76	40,700	14,746	
C18 reverse-phase	0.4	3.84	1.54	11,111	72,150	

Chromatography of the crude material over an immobilized bovine pancreatic kallikrein affinity column selectively isolated 0.013 % of the protein and 0.67 % of the trypsin inhibitory activity present. The majority of the trypsin inhibitory activity present in the starting supernatant did not bind to the immobilized kallikrein and is not related to bikunin (results not shown). Subsequent chromatography using C18 reverse-phase yielded a further purification of 5-fold, with a recovery of 0.2%. The final preparation was highly pure by SDS-PAGE (Figure 13), exhibiting an Mr of 21.3 kDa, and reacted on immunoblots to rabbit anti-placental bikunin 102-159 (not shown). N-terminal sequencing (26 cycles) yielded the expected sequence for mature placental bikunin (Figure 4F) starting at residue +1(ADRER....), showing that the senar peptide was correctly processed in the sequence.

alkylated, CNBr digested and then sequenced without resolution of the resulting fragments. Sequencing for 20 cycles yielded the following N-terminii:

	Sequence	Amount	Placental bikunin residue #
5	LRCFrQQENPP-PLG ADRERSIHDFCLVSKVVGRC FNYeEYCTANAVTGPCRASF PrY-V-dGS-Q-F-Y-G	21 pmol 20 pmol 16 pmol 6 pmol	154 - 168 (SEQ ID NO: 63) 1 - 20 (SEQ ID NO: 64) 100 - 119 (SEQ ID NO: 65) 25 - 43 (SEQ ID NO: 66)

Thus N-terminii corresponding to each of the expected four fragments were recovered. This confirms that the Sf9 expressed protein contained the entire ectodomain sequence of placental bikunin (1-170). N-terminal sequencing (50 cycles) of an additional sample of undigested Placental Bikunin (1-170) resulted in an amino acid sequence which at cycle 30 was devoid of any PTH-amino acid (PTH-asparagine was expected). A similar result was obtained upon sequencing of the natural protein from human placenta (Example 7) and is consistent with this residue being glycosylated as predicted from the amino acid sequence surrounding this asparagine residue. Furthermore, the cysteine residues within this region were also silent consistent with their participation in disulfide bonding.

### Example 10

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Inhibition specificity of purified placental bikunin derived from Sf9 cells.

The *in vitro* specificity of recombinant bikunin was determined using the materials and methods as described in Examples 3, 4 and 7. In addition, the inhibition of human tissue kallikrein by bikunin was measured by the incubation of 0.35 nM human tissue kallikrein recombinant bikunin in buffer containing 50 mM Tris (pH 9.0), 50 mM NaCl, and 0.01% triton x-100. After 5 min. at  $37^{\circ}$ C, 5  $\mu$ l of 2 mM PFR-AMC was added and the change in fluorescence monitored.

Inhibition of tissue plasminogen activator (tPA) was also determined as follows: tPA (single chain form from human melanoma cell culture from Sigma Chemical Co, St Louis, MO) was pre-incubated with inhibitor for 2 hr at room temperature in 20 mM Tris buffer pH 7.2 containing 150 mM NaCl, and 0.02% sodium azide. Reactions were subsequently initiated by transfer to a reaction system comprising the following initial component concentrations: tPA (7.5 nM), inhibitor 0 to 6.6 µM, DIle-Lpro-Larg-pNitroaniline (1mM) in 28 mM Tris buffer pH 8.5 containing 0.004 % (v/v) triton x-100 and 0.005% (v/v) sodium azide. Formation of p-Nitroaniline was determined from the A405nm measured following incubation at 37 C for 2hr.

The table below show the efficacy of recombinant bikunin as an inhibitor of various serine proteases in vitro. Data is shown compared against data obtained for

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screening inhibition using either recombinant bikunin, or aprotinin.

Table 9
Comparisons of Ki values for the inhibition of various proteases by recombinant placental bikunin (1-170) or aprotinin

Protease	Dana - Lieu - L	
(concentration)	Recombinant Bikunin Ki (nM)	Aprotinin Ki (nM)
Trypsin (48.5 pM)	0.064	0.8
Human Plasma Kallikrein (2.5 nM)	0.18	19.0
Human Tissue Kallikrein (0.35 nM)	0.04	0.004
Bovine Pancreatic Kallikrein (100 pM)	0.12	0.02
Human Plasmin (50 pM)	0.23	1.3
factor Xa (0.87 nM)	180	5% Inhibition at 31 μM
factor XIa (0.1 nM)	3.0	288
tissue plasminogen activator (7.5 nM)	< 60	no inhibition at 6.6 $\mu$ M
Tissue Factor VIIa	800	no inhibition at 1 μM

The results show that recombinant bikunin can be expressed in insect cells to yield an active protease inhibitor that is effective against at least five different serine protease inhibitors. Recombinant bikunin was more potent than aprotinin against human plasma kallikrein, trypsin and plasmin. Surprisingly, the recombinant bikunin was more potent that the synthetically derived bikunin fragments (7-64) and (102-159) against all enzymes tested. These data show that recombinant bikunin is more effective than aprotinin, using *in vitro* assays, and that one would expect better *in vivo* potency.

Besides measuring the potencies against specific proteases, the capacity of placental bikunin (1-170) to prolong the activated partial thromboplastin time (APTT) was evaluated and compared with the activity associated with aprotinin. Inhibitor was diluted in 20 mM Tris buffer pH 7.2 containing 150 mM NaCl and 0.02% sodium azide and added (0.1 ml) to a cuvette contained within an MLA Electra R 800 Automatic Coagulation Timer coagulometer (Medical Laboratory Automation Inc. Pleasantville N. V. The instrument was set to APTT and Sec. activation time and the dublicate mode mollowing addition of plasma (Specialty Assayed Reference Plasma tot 1-6-5185, Helena Laboratories, Beaumont, TX), the APTT reagent (Automated APTT-lot 102345, from Organon Teknika Corp., Durhan, NC) and 25 mM CaCl2 were automatically dispensed to inutiate clotting, and the clotting time was monitored automatically. The results

(Figure 14) showed that a doubling of the clotting time required approximately 2  $\mu$ M final aprotinin, but only 0.3  $\mu$ M Sf9 derived placental bikunin. These data show that placental bikunin is an effective anticoagulant, and useful as a medicament for diseases involving pathologic activation of the intrinsic pathway of coagulation.

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Although certain embodiments of the invention have been described in detail for the purpose of illustration, it will be readily apparent to those skilled in the art that the methods and formulations described herein may be modified without departing from the spirit and scope of the invention. Accordingly, the invention is not limited except as by the appended claims.

# WE CLAIM:

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1. A substantially purified protein, having serine protease inhibitory activity, selected from the group of proteins consisting of materials each of which comprises one of the following amino acid sequences, the amino acids of said sequences being numbered in accordance with the amino acid sequence of native human placental bikunin shown in figure 4F in which the N-terminal residue generated by removal of signal peptide is designated as residue 1:

10	ADRERSIHDF CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN	50
	YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRO DSEDHSSDMF	100
	NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE	150
	ACMLRCFRQQ ENPPLPLGSK	170
	(SEQ ID NO: 52);	
15		
	MAQLCGL RRSRAFLALL GSLLLSGVLA	-1
	Adrersiadf cluskuvgrc rasmprwwyn utdgscolfu yggcdgnsnn	50
	YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF	100
	NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE	150
20	ACMLRCFRQQ ENPPLPLGSK VVVLAGLFVM VLILFLGASM VYLIRVARRN	200
	QERALRIVWS SGDDKEQLVK NIYVL	225
	(SEQ ID NO: 49);	
	ADRERSIHDF CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN	50
25	YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF	100
	NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE	150
	ACMLRCFRQQ ENPPLPLGSK VVVLAGLFVM VLILFLGASM VYLIRVARRN	200
	QERALRIVWS SGDDKEQLVK NIYVL	225
	(SEQ ID NO: 70);	
30		
	AGSFLAWL GSLLLSGVLA -1	
	LOBERCARTE CAMPANALEC BY CHERMANA CHARLES EL COULTANONA	
	TKEECLE: "TTENAT Law ISRNAAL SAKER LEGENDE	
25	MYEEYCTANA VTGPCRASEP RWYEDVERNS NNFIYGGCE GNKNSYRSEE	
35	ACMLRCFRQQ ENPPLPLGSK VVVLAGAVS	179
	(SEQ ID NO: 2);	

	MLR AEADGVSRLL GSLLLSGVL	-A -1
	ADRERSIHDF CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN	50
	YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF	100
	NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE	150
5	ACMLRCFRQQ ENPPLPLGSK VVVLAGLFVM VLILFLGASM VYLIRVARRN	200
	QERALRTVWS SGDDKEQLVK NTYVL	225
	(SEQ ID NO: 45);	
	MAQLCGL RRSRAFLALL GSLLLSGVLA	<b>A</b> -1
10	ADRERSIHDF CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN	50
	YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF	100
	NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE	150
	ACMLRCFRQQ ENPPLPLGSK VVVLAGLFVM VLILFLGASM VYLIRVARRN	200
	QERALRIVWS FGD	213
15	(SEQ ID NO: 47);	
	ADRERSIHDF CLVSKVVGRC RASHPRWWYN VTDGSCQLFV YGGCDGNSNN	50
	YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF	100
	NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE	150
20	ACMLRCFRQQ ENPPLPLGSK VVVLAGLFVM VLILFLGASM VYLIRVARRN	200
	QERALRIVWS FGD	213
	(SEQ ID NO: 71);	
	IHDF CLVSKVVGRC RASHPRWWYN VTDGSCQLFV YGGCDGNSNN 5	0
25	YLTKEECLKK CATV 6.	4
	(SEQ ID NO: 4);	
	CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN 50	_
	YLTREECLKK C	1
30	(SEQ ID NO: 5);	
	YEEYCTANA VTGPCRASFP RWYFDVERNS CHNFIYGGCR GNKNSYRSEE 15	
	ACMLRCFRQ	159
	(SEQ ID NO: 6);	
35		••
	CTANAVIGPC RASFPRWYFD VERNSCHNFI YGGCRGNKNS YRSEE 15	
	ACMLRC 15	56

	(SEQ ID NO: 7);	
	IHDF CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN	50
	YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF	75
5	NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE	125
	ACMLRCFRQ	159
	(SEQ ID NO: 3);	
	CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN	50
10	YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF	100
	NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE	150
	ACMLRC	156
	(SEQ ID NO: 50);	
15	ADRERSIHDF CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSMN	25
	YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF	75
	NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE	125
	ACMLRCFRQQ ENPPLPLGSK VVVLAGAVS	179
	(SEQ ID NO: 1); and	
20		
	ADRERSIHDF CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN	50
	YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DS	2
	(SEQ ID NO: 8).	

- 25 2. A protein as in claim 1, wherein said protein is glycosylated, or contains at least one intra-chain cysteine-cysteine disulfide bond, or is both glycosylated and contains at least one intra-chain cysteine-cysteine disulfide bond.
- 3. A pharmaceutical composition for inhibiting serine protease activity, comprising a protein of claim 1 or claim 2 plus a pharmaceutically acceptable carrier.
- 35 5. A self-replicating protein expression vector containing a nucleic acid sequence which encodes for and is capable of expressing a protein of claim or claim 2.

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A method for inhibiting serine protease activity comprising contacting 6. serine protease with an effective amount of at least one protein of claim 1 or claim 2.

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A method for treating a condition of brain edema, spinal cord edema, 7. multiple sclerosis, ischemia, perioperative blood loss, sepsis, septic shock, fibrosis, disease associated with pathologic blood coagulation or clotting, polytrauma, stroke, cerebral or subarachnoid hemorrhage, inflamation of the brain, inflamation of the spinal cord, cerebral infection, cerebral granulomatosis, spinal infection, spinal granulomatosis, open heart surgery, gastric cancer, cervical cancer, or prevention of metastasis comprising administering to a subject having such a condition an effective amount of the protein of claim 1 or claim 2.

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The method of Claim 7 wherein said condition is brain edema, spinal 8. cord edema, multiple sclerosis, ischemia, perioperative blood loss, sepsis, septic shock, fibrosis, disease associated with pathologic blood coagulation or clotting, stroke, cerebral or subarachnoid hemorrhage, inflamation of the brain, inflamation of the spinal cord, cerebral infection, cerebral granulomatosis, spinal infection, spinal granulomatosis, or open heart surgery.

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The method of Claim 7 wherein said condition is gastric cancer, cervical 9. cancer, or prevention of metastasis. A method for the preparation of a medicament for the treatment of brain

edema, spinal cord edema, multiple sclerosis, ischemia, perioperative blood loss, sepsis, septic shock, fibrosis, disease associated with pathologic blood coagulation or clotting, stroke, cerebral or subarachnoid hemorrhage, inflamation of the brain, inflamation of the spinal cord, cerebral infection, 30 cerebral granulomatosis, spinal infection, spinal granulomatosis, open heart surgery, gastric cancer, cervical cancer, or prevention of metastasis.

A method for preparing a protien of claim 1 or claim 2 using 11. recombinant DNA technology.

# FIGURE 1

	GCCCGGGTCG	TTTCTCGCCT	GGCTGGGATC	GCTGCTCCTC	TCTGGGGTCC	50
ORF	PGR	F S P	G W D R	c s s	L G S	16
R35464	TGGCCGGCCG	ACCGAGAACG	CAGCATCCAC	GACTTCTGCC	TGGTGTCGAA	100
ORF	M S Y D		S I H	D F C L		33
R35464	GGTGGTGGGC	AGATTCCGGG	CCTCCATGCC	TAGGTGGTGG	TACAATGTCA	150
ORF		RERA		R W W	Y N V T	50
R35464	CTGACGGATC	CTGCCAGCTG	TTTGTGTATG	GGGGCTGTGA	CGGAAACAGC	200
ORF		CQL			G N S	66
	AATAATTACC	TGACCAAGGA	GGAGTGCCTC	AAGAAATGTG	CCACTGTCAC	250
ORF	ииуг	T K E	E C L	K K C A	T V T	83
	AGAGAATGCC	ACGGGTGACC	TGGCCACCAG	CAGGAATGCA	GCGGATTCCT	300
ORF	ENA	TGDL	A T S	RNA	A D S S	100
		TGCTCCCAGA		CTTGAAGACC	ACTTCAGCGA	350
ORF	V P S	A P R	R Q D S	• R P	L Q R	116
		NTATTGNAAG		CCGNCAACGN	ATT	393
ORF	Y V S *	I * A	IIA	P • T •		130
KEY						
R35464	- Nucleic	cid sequenc	e_of EST R3	5464 (SEQ I	D NO: 12)	

ORF = EST R35464 Open Reading Frame Translation (SEQ ID NO: 13)

# FIGURE 2

R74593 ORF	Q · L			Q E M	C H C H	50 17
R74593 ORF			CCTGGCCACC P G H Q	AGCAGGAATG Q E C	CAGCGGATTC S G F	100 33
			AAGGCAGGAT R Q D	TCTGAAGACC S E D H	ACTCCAGCGA S S D	150 50
	TATGTTCAAC M F N		ACTGCACCGC C T A	CAACGCAGTC N A V	ACTGGGCCTT T G P C	200 67
R74593 ORF	GCCGTGCATC R A S		TGGTACTTTG W Y F D	ACGTGGAGAG V E R	GAACTCCTGC N S C	250 83
R74593 O <b>RF</b>			CTGCCGGGGC C R G	AATAAGAACA N K N S	GCTACCGCTC Y R S	300 100
R74593 O <b>RF</b>	TGAGGAGGCC E E A	TGCATGCTCC C M L R		CCAGCAGGAG Q Q E	AATCCTCCCC N P P L	350 117
R74593 O <b>RF</b>	TGCCCCTTGG P L G		GTGGTTCTGG V V L A	CCGGGGCTGT G A V	TTCGTGATGG S * W	400 133
R74593 O <b>RF</b>	TGTTGATCCT C * S F			GTCTTACTGA V L L I	TTCCGGGTGG P G G	450 150
R74593 O <b>RF</b>	CAAGGAGGAA K E E		GCCCTGCGGA P A X	ncgtctggag R L E	CTTCGGAGAT L R R *	500 167
	GACAAGGGNT O G					510

#### KEY

R74593 - Nucleic acid sequence of EST R74593 (SEQ ID NO: 14)
ORF - EST R74593 Open Reading Frame Translation (SEQ ID NO: 15)

### FIGURE 3

R35464 N39798	GGC	CGG	GICGI	TTC	TCG	CCTG			A-TC Antc								50 23
H94519	GC	NGC	G-CGT	TNN	TCG	CNT-											47
R74593 corr.										301				<b>G</b>	301	CG	4 /
Consensus	GGC	caa	GTCGI		TCC	CCTG	CCT	ccc	1 - TC	CCT							5.5
Translation	A	G	S F				L	G	<u>A</u> -≀C	L						در	50
1141131401011	^	J	<b>3</b> F		^			G	3	L	L	L	S	G	٧		- 3
R35464	TC	300	GGCCG	. ACC	GAG	AACG	CAG	CAT	CCAC	GAC	**~	*~~~	*~~	~~			100
N39798			GGCCG														100
H94519			GGCCG														96
R74593 corr.										onc.			100	101		~~	75
Consensus	TG	G-C	GGCCG	ACC	GAG	AACG	CAG	CAT	CCAC	GAC	TTC	TGCC	tcc	TG	TCG	 A A	99
Translation	L		A D			R	S		H	ם						r. K	15
	_			•	_	-	-	•	-	_	-			•	•	2	
R35464	GG:	rgg	TGGGC	AGA	TTC	CGGG	COT	CCA'	TGCC	TAG	GTG	GTGG	TAC	331	CT	٠,	150
N39798			TGGGC														
H94519			TGGGC														
R74593 corr.																	110
Consensus	GG1	rgg <sup>,</sup>	TGGGC	AGA	TGC	CGGG	CCT	CCA	TGCC	TAG	GTG	GTGG	TAC	AAI	CGT	CA	149
Translation	¥				_	R A	s			R	Ħ	H	Y		Y	ī	32
							_	_	_	_	_	-	•	••	-		7.
R35464	CTC	SAC	GGATC	CTG	CCA	GCTG	TTT	STG	TATG	GGG	GCT	GTGA	CGG		CAC	30	200
N39798	CTC	SAC	GGATC	CTG	CCA	GCTG	TTT	GTG	TATG	GGG	GCT	STGA	CGG	222	CAC	30	177
H94519	CTO	AC	GGATC	CTG	CA	GCTG	TIT	GTG	PATG	GGG	GCT	GTGA	CGG	AAA	CAC	30	196
R74593 corr.															(		2
Consensus	CTC	AC	GGATC	CTG	CA	GCTG	TTT	GTG	CATG	GGG	GCT	STGA	CGG				
Translation	1		<u>a</u> s	C		L	Ε :				C				2	•	48
												_		_	-		-
R35464			PTACC														
R35464 N39798	AAT	: 1	PTACC	TGA	CA	AGGA	GGA	STGO	CTC	AAG	AAA:	TGTG	CCA	CTG	TCJ	\C	227
N39798 H94519	AAI	'AA'	CTACC	TGA(	CA	AGGA AGGA	GGA(	STG(	CTC	AAG	NAA:	rgtg rgtg	CCA	CTG	TC	IC IC	227
N39798	AAI	'AA'	COATT COATT	TGA( TGA( TGA(	CA CA	AGGA AGGA AGGA	GGA( GGA( GGA(	STGC STGC	CTC	AAGI AAGI AAGI	NAA! NAA! NAA!	rgtg rgtg rgtg	CCA	CTG CTG	TCJ TCJ	ic ic	227 246 52
N39798 H94519 R74593 corr. Consensus	AAT	\ \ \ \ \ \	COATT COATT	TGA( TGA( TGA(	CA CA	AGGA AGGA	GGA( GGA( GGA(	STGC STGC	CTC	AAGI AAGI AAGI	NAA! NAA! NAA!	rgtg rgtg rgtg	CCA	CTG CTG	TCJ TCJ	ic ic	227 246 52
N39798 H94519 R74593 corr.	AAT	\ \ \ \ \ \	COATT COATT	TGA( TGA( TGA(	CA CA	AGGA AGGA AGGA AGGA	GGA( GGA( GGA(	STGC STGC	CTC	AAGI AAGI AAGI	WAS WAS WAS	rgtg rgtg rgtg	CCA	CTG CTG CTG	TCI TCI TCI	ic ic	227 246 52
N39798 H94519 R74593 corr. Consensus Translation	AAT AAT AAT B	iaai iaai iaai iaai	TACC TACC TACC TACC	TGA( TGA( TGA( TGA(	CAL CAL CAL CAL K	AGGA AGGA AGGA AGGA E	GGA( GGA( GGA( GGA(	STGC STGC STGC C	CTC CTC CTC L	AAGI AAGI AAGI K	NAA! NAA! NAA! NAA!	rgtg rgtg rgtg rgtg C A	CCA CCA CCA CCA	ete ete ete ete	TCJ TCJ TCJ	ic ic ic	227 246 52 249 65
N39798 H94519 R74593 corr. Consensus Translation	AAT AAT AAT N	AATAATAATAATAATAATAATAATAATAATAATAATAAT	PTACC PTACC PTACC TTACC Y L	TGAG TGAG TGAG TGAG	CCAL CCAL CCAL CCAL K	AGGA AGGA AGGA E BACC	GGAG GGAG GGAG E	etgo etgo etgo c c	CTC CTC CTC L	AAGI AAGI AAGI K	NAA! NAA! NAA! K	rgtg rgtg rgtg rgtg C A	CCA CCA CCA CCA	CTG CTG CTG	TCJ TCJ TCJ TCJ	ic ic ic ic it	227 246 52 249 65
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798	AAT AAT AAT N AGA	AAT AAT AAT B B GAJ	PTACC PTACC PTACC TTACC Y L ATGCC ATGCC	TGAG TGAG TGAG TGAG ACGG	CCAL CCAL CCAL CCAL CCAL CCAL CCAL CCAL	AGGA AGGA AGGA E BACC GACC	GGAG GGAG GGAG E TGGG	ETGO ETGO ETGO C C CAO	CTC CTC CTC L CCAG	AAGI AAGI AAGI K CAGC	NAA! NAA! NAA! K SAA!	rete rete rete rete c a reca reca	CCA CCA CCA CCA T	CTG CTG CTG CTG CTG	TCJ TCJ TCJ		227 246 52 249 65 300 277
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519	AAT AAT AAT N AGA AGA	AAT AAT AAT B GAJ GAJ	PTACC PTACC PTACC PTACC Y L ATGCC ATGCC	TGAG TGAG TGAG TGAG ACGG ACGG	CCAL CCAL CCAL CCAL K GGT( GGT(	AGGA AGGA AGGA E BACC BACC BACC	GGAG GGAG GGAG E TGGG TGGG	ETGO ETGO ETGO C CAO CAO CAO	CTC CTC CTC L CAG CAG CAG	AAGI AAGI AAGI K CAGC CAGC	NAA! NAA! NAA! K SAA! SAA!	rete rete rete rete a reca reca reca	CCA CCA CCA CCA CCA CCA CCA CCA CCA CCA	CTG CTG CTG CTG CTG CTG	TCJ TCJ TCJ TCG		227 246 52 249 65 300 277 296
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798	AAT AAT AAT N AGA AGA AGA	IAAI IAAI IAAI B IGAI IGAI IGAI	PTACE PTACE PTACE Y L  ATGCE ATGCE ATGCE ATGCE	TGAG TGAG TGAG TGAG ACGG ACGG ACGG	CCAL CCAL CCAL CCAL CCAL CCAL CCAL CCAL	AGGA AGGA AGGA E BACC BACC BACC BACC	GGAG GGAG GGAG E TGGG TGGG TGGG	etgo etgo etgo c c c c c c c c	ICTC ICTC ICTC ICTC ICAG ICAG ICAG	AAGI AAGI AAGI K CAGI CAGI CAGI CAGI	NAA! NAA! NAA! K SAA! SAA!	rete rete rete rete . A reca reca reca	CCAI CCAI CCAI CCAI T GCGG GCGG GCGG	CTG CTG CTG CTG GAT GAT GAT	TCI TCI TCI TCI TCI		227 246 52 249 65 300 277 296 102
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr. Consensus	AAT AAT AAT N AGA AGA AGA	IAAI IAAI IAAI IGAU IGAU IGAU IGAU	PTACC PTACC PTACC PTACC Y L NTGCC NTGCC NTGCC NTGCC NTGCC	TGAG TGAG TGAG TGAG ACGG ACGG ACGG ACGG	CAL CAL CAL CAL CAL CAL CAL CAL CAL CAL	AGGA AGGA AGGA E BACC BACC BACC BACC BACC	GGAG GGAG GGAG TGGG TGGG TGGG	STGC STGC STGC STGC SCAC SCAC SCAC	COTO COTO L COAG COAG COAG COAG COAG	AAGI AAGI AAGI AAGI CAGC CAGC CAGC	NAA! NAA! NAA! K SAA! SAA! SAA!	rete rete rete rete a reca reca reca reca	CCA CCA CCA CCA CCA CCA CCA GCG GCG GCG	CTG CTG CTG CTG CTG SAT GAT GAT	TC0 TC0 TC0 TC0 TC0		227 246 52 249 65 300 277 296 102
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr.	AAT AAT AAT N AGA AGA AGA	IAAI IAAI IAAI IGAU IGAU IGAU IGAU	PTACE PTACE PTACE Y L  ATGCE ATGCE ATGCE ATGCE	TGAG TGAG TGAG TGAG ACGG ACGG ACGG ACGG	CAL CAL CAL CAL CAL CAL CAL CAL CAL CAL	AGGA AGGA AGGA E BACC BACC BACC BACC	GGAG GGAG GGAG TGGG TGGG TGGG	STGC STGC STGC STGC SCAC SCAC SCAC	COTO COTO L COAG COAG COAG COAG COAG	AAGI AAGI AAGI AAGI CAGC CAGC CAGC	NAA! NAA! NAA! K SAA! SAA! SAA!	rete rete rete rete a reca reca reca reca	CCA CCA CCA CCA CCA CCA CCA GCG GCG GCG	CTG CTG CTG CTG CTG SAT GAT GAT	TC0 TC0 TC0 TC0 TC0		227 246 52 249 65 300 277 296 102
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr. Consensus Translation	AAT AAT AAT N AGA AGA AGA AGA	IAAT IAAT IAAT IGAU IGAU IGAU IGAU IGAU	TTACC TTACC TTACC TTACC TTACC TTACC TTACC Y L  ATGCC ATGCC ATGCC A	TGAG TGAG TGAG TGAG ACGG ACGG ACGG ACGG	CAL CAL CAL SGTO SGTO SGTO SGTO	AGGA AGGA AGGA E SACC SACC SACC SACC SACC	GGAG GGAG GGAG E TGGG TGGG TGGG	STGC STGC STGC SCAC SCAC SCAC SCAC	ICTC ICTC ICTC L ICAG ICAG ICAG ICAG ICAG ICAG ICAG	AAGI AAGI AAGI K CAGC CAGC CAGC CAGC	AAA: AAA: AAA: K SAA: SAA: SAA:	rete rete rete rete c a reca reca reca reca	CCAI CCAI CCAI CCAI CCAI GCGI GCGI GCGI	CTG CTG CTG CTG CTG CAT GAT GAT GAT	TCO TCO TCO TCO TCO TCO	IC IC IC IT	227 246 52 249 65 300 277 296 102 299 82
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr. Consensus Translation	AAT AAT AAT N AGA AGA AGA AGA AGA	TAAT TAAT TAAT IGAU IGAU IGAU IGAU IGAU	TTACC	TGAG TGAG TGAG TGAG ACGC ACGC ACGC ACGC	CCAL CCAL CCAL CCAL CCAL CCAL CCAL CCAL	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAG GGAG GGAG E TGGG TGGG TGGG A	STGC STGC STGC STGC SCAC SCAC SCAC T	ICTC ICTC ICTC ICAG ICAG ICAG ICAG ICAG ICAG ICAG ICA	AAGI AAGI AAGI K CAGC CAGC CAGC CAGC	MAA: MAA: MAA: K SAA: SAA: SAA: SAA:	rete rete rete rete c a reca reca reca reca a	CCAI CCAI CCAI CCAI CCAI GCGG GCGG GCGG	CTG CTG CTG CTG CTG SAT GAT GAT GAT	TCJ TCJ TCJ TCG TCG TCG	IC IC IC IT IT IT IT IS GA	227 246 52 249 65 300 277 296 102 299 82
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr. Consensus Translation R35464 N39798	AAT AAT AAT N AGA AGA AGA AGA CTC	TAAT TAAT TAAT TAAT GAU GGU GGU GGU TTCC TTCC	TTACC TTACC TTACC TTACC TTACC TTACC Y L  ATGCC ATGCC ATGCC A  CCAAG CCAAG	TGAG TGAG TGAG TGAG ACGC ACGC ACGC ACGC	CALCALICATION OF THE COLOR OF T	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAG GGAG GGAG E TGGG TGGG TGGG TGGG A AGGG	STOCK	CCTC CCTC L CCAG CCAG CCAG CCAG CCAG CCAG CCAG C	AAGI AAGI AAGI K CAGC CAGC CAGC CAGC CAGC CAGC	AAA AAA K K GAA GAA GAA	rete rete rete reca reca reca reca reca a	CCAI CCAI CCAI CCAI CCAI GCGG GCGG GCGG	CTG CTG CTG CTG CTG SAT SAT SAT SAT SAT SAT SAT SAT SAT SAT	TCI TCI TCI TCI TCI TCI TCI TCI TCI TCI	IC IC IC IT IT IT IT IS GA	227 246 52 249 65 300 277 296 102 299 82 350 326
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519	AAT AAT AAT AGA AGA AGA AGA CTC	TAAT TAAT TAAT NAAT NGAU NGGU NGGU NGGU TCC	TTACC TTACC TTACC TTACC TTACC TTACC Y L  ATGCC ATGCC ATGCC A  CCAAG CCAAG CCAAG	TGAG TGAG TGAG TGAG ACGC ACGC ACGC ACGC	CALCALLANDER OF THE COLORS OF	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAGGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGG	STAGE	CCTC CCTC L CCAG CCAG CCAG CCAG CCAG CCAG CCAG C	AAGI AAGI AAGI K CAGC CAGC CAGC CAGC CAGC CAGC CAGC CA	AAA: AAA: AAA: K ( GGAA: GGAA: GGAA: GGAA:	rete rete rete reca reca reca reca reca a reca sacc sacc	CCAI CCAI CCAI CCAI GCGG GCGG GCGG GCGG	CTG CTG CTG CTG CTG SAT SAT SAT SAT SAT SAT SAT SAT	TCU	COLOR TITTES GAMA	227 246 52 249 65 300 277 296 102 299 82 350 326 345
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr.	AAT AAT AAT AGA AGA AGA AGA CTC CTC	TAAT TAAT TAAT TAAT TAAT GGAJ GGAJ GGAJ	ITACO	TGAG TGAG TGAG TGAG TGAG ACGC ACGC ACGC	COOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOC	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAGGGAGGGAGGGAGGGAGGGAGGGAGGGAGGGAGGG	STOCE	CCTC CCTC L CCAG CCAG CCAG CCAG CCAG CCAG CCAG C	AAGI AAGI AAGI AAGI CAGC CAGC CAGC CAGC	AAA: AAA: AAA: K ( ) GAA: GAA: GAA: GAA: GAA: GAA:	rete rete rete reca reca reca reca reca a sacc sacc sacc	CCAI CCAI CCAI CCAI GCGG GCGG GCGG GCGG	CTG CTG CTG CTG CTG GAT GAT GAT CCA CCA	TCU TCU TCU TCU TCU TCU TCU TCU TCU TCU	COLOR TITTES GARAGA	227 246 52 249 65 300 277 296 102 299 82 350 326 345 151
N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Consensus Consensus	AAT AAT AAT AGA AGA AGA AGA CTG CTG CTG	IAAI IAAI IAAI IGAU IGAU IGGU ITCC ITCC ITCC	ITACC	TGAC TGAC TGAC ACGC ACGC ACGC ACGC TGCT TGCT	CALCOLOR CONTINUES CONTINU	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAG GGAG GGAG E TGGG TGGG TGGG TGGG AGGG A	STOCKER TO THE STOCKE	CCTC CCTC L CCAG CCAG CCAG CCAG CCAG CCAG CCAG C	AAGI AAGI AAGI K CAGC CAGC CAGC CAGC CAGC CAGC CAGC CA	AAA: AAA: AAA: K GAA: GAA: GAA: GAA: GAA	rete rete rete reca reca reca reca reca reca reca rec	GCAI CCAI CCAI CCAI CCAI CCAI GCGG GCGG	CTG CTG CTG CTG CTG GAT GAT GAT CCA CCA CCA	TCU TCU TCU TCU TCU TCU TCU TCU TCU TCU	COLOR TITTES GARAGA	227 246 52 249 65 300 277 296 102 299 82 350 326 345 151 348
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr.	AAT AAT AAT AGA AGA AGA AGA CTG CTG CTG	IAAI IAAI IAAI IGAU IGAU IGGU ITCC ITCC ITCC	ITACO	TGAC TGAC TGAC ACGC ACGC ACGC ACGC TGCT TGCT	CALCOLOR CONTINUES CONTINU	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAG GGAG GGAG E TGGG TGGG TGGG TGGG AGGG A	STOCKER TO THE STOCKE	CCTC CCTC L CCAG CCAG CCAG CCAG CCAG CCAG CCAG C	AAGI AAGI AAGI K CAGC CAGC CAGC CAGC CAGC CAGC CAGC CA	AAA: AAA: AAA: K GAA: GAA: GAA: GAA: GAA	rete rete rete reca reca reca reca reca a sacc sacc sacc	GCAI CCAI CCAI CCAI CCAI CCAI GCGG GCGG	CTG CTG CTG CTG CTG GAT GAT GAT CCA CCA CCA	TCU TCU TCU TCU TCU TCU TCU TCU TCU TCU	COLOR TITTES GARAGA	227 246 52 249 65 300 277 296 102 299 82 350 326 345 151
N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Translation	AAT AAT AAT AGA AGA AGA AGA CTG CTG CTG	I AAT I AAT	TTACC TTACC TTACC TTACC TTACC TTACC Y L  ATGCC ATGCC ATGCC A  CCAAG CCAAG CCAAG CCAAG CCAAG CCAAG	TGAG TGAG TGAG TGAG TGAG ACGC ACGC ACGC	COAL COAL ECAL EGGIO EGGIO EGGIO EGGIO EGGIO EGGIO EGGIO EGGIO EGGI EGGI	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAGGGAGGGAGGGAGGGAGGGAGGGAGGGAGGGAGGG	STGC STGC STGC STGC SCAC SCAC SCAC SCAC SCAC SCAC SCAC SC	CCTC CCTC L CCAG CCAG CCAG CCAG CCAG CCAG CCAG C	AAGI AAGI AAGI AAGI K CAGC CAGC CAGC CAGC CAGC CAGC CAGC CA	AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	TGTG TGTG TGTG TGCA TGCA TGCA TGCA TGCA	GCAI CCAI CCAI CCAI CCAI CCAI GCGG GCGG	CTG CTG CTG CTG CTG GAT GAT GAT CCA CCA CCA	TCU TCU TCU TCU TCU TCU TCU TCU TCU TCU	COLOR TITTES GARAGA	227 246 52 249 65 300 277 296 102 299 82 350 326 345 151 348
N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Translation	AAT AAT AAT N AGA AGA AGA AGA CTG CTG	I AAT TOO TOO TOO TOO TOO TOO TOO TOO TOO	TTACC	TGAG TGAG TGAG TGAG ACGC ACGC ACGC ACGC	CALCOLOR P	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAGGGAGGGAGGGAGGGAGGGAGGGAGGGAGGGAGGG	STOCKER TO THE STOCKE	CCTC CCTC CCTC L CCAG CCAG CCAG CCAG CCAG CCAG CCAG C	AAGI AAGI AAGI CAGC CAGC CAGC CAGC CAGC	AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	IGTG IGTG IGTG IGCA IGCA IGCA IGCA IGCA IGCA IGCA IGC	GCAI CCAI CCAI CCAI GCGI GCGI GCGI GCGI	CTG CTG CTG CTG CTG SAT GAT GAT GAT CCA CCA SCCA S	TCU TCU TCU TCU TCU TCU TCU TCU TCU TCU	COLOR TITTETS GARAGA	227 246 52 249 65 300 277 296 102 299 82 350 326 345 151 348
N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Translation	AAT AAT NA AGA AGA AGA AGA AGA AGA AGA AGA AGA	I AAI I AAI I AAI I GAU I GAU I GAU I GAU I TOO I TOO	TTACC TTGCC	TGAG TGAG TGAG TGAG ACGC ACGC ACGC ACGC	COMPANY OF THE PROPERTY OF THE	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAGGGAGGGAGGGAGGGAGGGAGGGAGGGAGGGAGGG	STOCKER TO THE STOCKE	CCTC CCTC CCAG CCAG CCAG CCAG CCAG CCAG	AAGI AAGI AAGI K CAGC CAGC CAGC CAGC CAGC CAGC CAGC CA	AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	IGTG IGTG IGTG IGCA IGCA IGCA IGCA IGCA IGCA IGCA IGC	GCAI CCAI CCAI CCAI GCGI GCGI GCGI GCGI	CTG CTG CTG CTG CTG GAT GAT GAT GAT CCA CCA S	TOCO TOCO TOCO TOCO TOCO TOCO TOCO TOCO	COLOR TITITES GARAGA	227 246 52 249 65 300 277 296 102 299 82 350 326 345 151 348 98
N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Translation	AAT	IAAN IAAN IAAN IAAN IAAN IAAN IAAN IAAN	TTACC	TGAG TGAG TGAG ACGC ACGC ACGC ACGC TGCT TGCT	COAL COAL ECAL EGGICAGE EGGICA EGGICA EGGICA EGGICA EGGIC	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAGGGAGGGAGGGAGGGAAGGGAAGGGAAGGGAAGGGAAGGGAAGGGAAGGGAAGGGAAGGAAATAAATA	STOCKER TO THE PROPERTY OF THE	CCTC CCTC CCTC CCAG CCAG CCAG CCAG CCAG	AAGI AAGI AAGI CAGC CAGC CAGC CAGC CAGC	AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	rete rete rete rete rete rete rete rete	GCAI CCAI CCAI CCAI GCGI GCGI GCGI GCGI	CTG CTG CTG CTG CTG GAT GAT GAT GAT CCA CCA CCA CCA	TOCO TOCO TOCO TOCO TOCO TOCO TOCO TOCO	COUNTY TITTE SALAMAN GG	227 246 52 249 65 300 277 296 102 299 82 350 326 345 151 348 98
N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr. Consensus Translation R35464 N39798 H94519 R74593 corr. Consensus Translation	AAT	I AAII I AAII	TTACC TTGCC	TGAG TGAG TGAG TGAG ACGC ACGC ACGC ACGC	CCAL CCAL ECCAL EC	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAGGGAGGGAGGGAGGGAAATAAAA	STGC STGC STGC STGC STGC STGC STGC STGC	CCTC CCTC CCTC CCAG CCAG CCAG CCAG CCAG	AAGI AAGI AAGI CAGC CAGC CAGC CAGC CAGC	AAA: AAA: AAA: AAA: AAA: AAA: AAA: AAA	IGTG IGTG IGTG IGTG IGCA IGCA IGCA IGCA IGCA IGCA IGCA IGC	GCAI CCAI CCAI CCAI GCGI GCGI GCGI GCGI	CTGCCTGCCTGCCTGCCTGCCTGCCTGCCTGCCTGCCTG	TOCO TOCO TOCO TOCO TOCO TOCO TOCO TOCO	COUNT TITTE SALAMA COOK	227 246 52 249 65 300 277 296 102 299 82 350 326 345 151 348 98
N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Translation  R35464 N39798 H94519 R74593 corr. Consensus Translation	AAT	GAN	TTACC	TGAG TGAG TGAG TGAG ACGC ACGC ACGC ACGC	CCAL CCAL GGT GGT GGT GGT CCC CCCC CCCC CCCC CCC	AGGA AGGA AGGA AGGA E BACC BACC BACC BACC BACC BACC BACC B	GGAGGGAGGGAGGGAGGGAAATAAAA	STGC STGC STGC STGC STGC STGC STGC STGC	CCTC CCTC CCTC CCAG CCAG CCAG CCAG CCAG	AAGI AAGI AAGI CAGC CAGC CAGC CAGC CAGC	ANA: ANA: ANA: ANA: ANA: ANA: ANA: ANA:	IGTG IGTG IGTG IGTG IGCA IGCA IGCA IGCA IGCA IGCA IGCA IGC	GCAI CCAI CCAI CCAI GCGI GCGI GCGI GCGI	CTGCTGCTGGATGATGATGATGATGATGATGATGATGATGATGATGA	TOCOTOCO SCOOL TOCOTOCO TOCOTOCO TOCOTOCO TOCOTOCO TOCOTOCO	COCO TITITIES GARAGA COCOGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	227 246 52 249 65 300 277 296 102 299 82 350 326 345 151 348 98

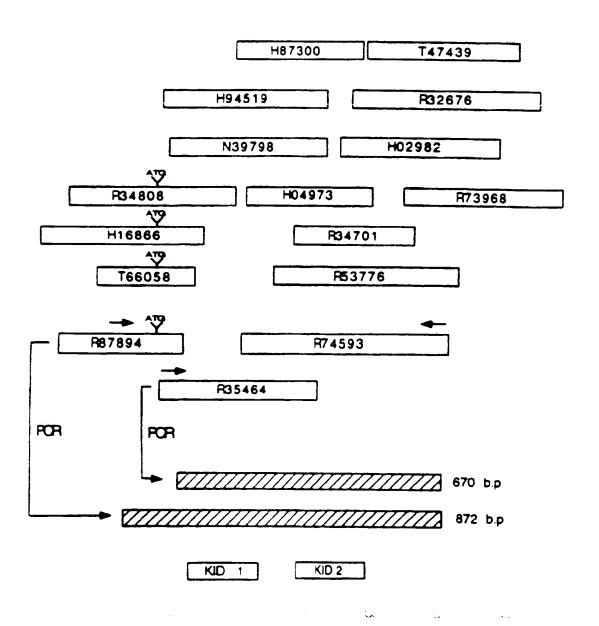
#### FIGURE 3 (CONT)

R35464		
<b>н39798</b>	CCTTGC-GTG GAATCCTTTC CCACGCTGGN AATTTNGACG TTGAGAAGGA 421	
H94519	CCT-GC-GTG -CATCCTT-C CCACGCTGGT ACTTT-GNCG 427	
R74593 corr.	CCTTGCCGTG -CATCCTT-C CCACGCTGGT ACTTT-GACG TGGAGA-GGA 243	
Consensus	CCTTGCCGTG -CATCCTT-C CCACGCTGGT ACTTT-GACG TGGAGA-GGA 440	
Translation	PCRASEPRWYFDVERN129	
	TO A A DI I WATER DV Z R N 129	
R35464	**********	
	AC 423	
N39798	423	
H94519		
R74593 corr.	ACTCCTGCAA TAACTTCATC TATGGAGGCT GCCGGGGCAA TAAGAACAGC 293	
Consensus	ACTCCTGCAA TAACTTCATC TATGGAGGCT GCCGGGGCAA TAAGAACAGC 490	
Translation	SCN NFI Y GGC RGN KNS 145	
R35464		
N39798	***************************************	
H94519		
R74593 corr.	TACCGCTCTG AGGAGGCCTG CATGCTCCGC TGCTTCCGCC AGCAGGAGAA 343	
Consensus	TACCGCTCTG AGGAGGCCTG CATGCTCCGC TGCTTCCGCC AGCAGGAGAA 540	
Translation	YRSEEACHLRCFRQQEN 162	
R35464		
N39798		
H94519		
R74593 corr.	TCCTCCCCTG CCCCTTGGCT CAAAGGTGGT GGTTCTGGCC GGGGCTGTTT 393	
Consensus	TCCTCCCCTG CCCCTTGGCT CAAAGGTGGT GGTTCTGGCC GGGGCTGTTT 590	
Translation	P P L P L G S K V V V L A G A V S 179	
	T T T T T T T T T T T T T T T T T T T	
R35464		
N39798	******************************	
H94519		
R74593 corr.	CGTGATGGTG TTGATCCTTT TCCTGGGGAG CNTCCATGGT CTTACTGATT 443	
Consensus	CGTGATGGTG TTGATCCTTT TCCTGGGGAG CNTCCATGGT CTTACTGATT 640	
Translation	* W C * S F S W G A S M V L L I 195	
-20104		
R35464		
N39798	***************************************	
H94519		
R74593 corr.	CCGGGTGGCA AGGAGGAACC AGGAGCGTGC CCTGCGGANC GTCTGGAGCT 493	
Consensus	CCGGGTGGCA AGGAGGAACC AGGAGCGTGC CCTGCGGANC GTCTGGAGCT 690	
Translation	PGGKEEPGACPA*RLEL212	
R35464		
N39798	••••••	
H94519		
R74593 corr.	TCGGAGATGA CAAGGGNT 511	
Consensus	TCGGAGATGA CAAGGGNT 708	
Translation		
	R R * Q G 217	
KEY		
	Aid said samenas of PCP D26464 (ABS to US . 15)	
M30700 - MUC1(	eic acid sequence of EST R35464 (SEQ ID NO.: 12)	
MISTA - MACTO	eic acid sequence of EST N39798 (SEQ ID NO.: 17)	

H94519 - Nucleic acid sequence of EST H94519 (SEQ ID NO.: 16) R74593 corr. - Corrected version of (SEQ ID NO.: 14) G at b.p. 114 Consensus - Nucelic acid sequence for human bikunin (SEQ ID NO.: 9) Translation - Amino acid Translation of Consensus (SEQ ID NO.: 10)

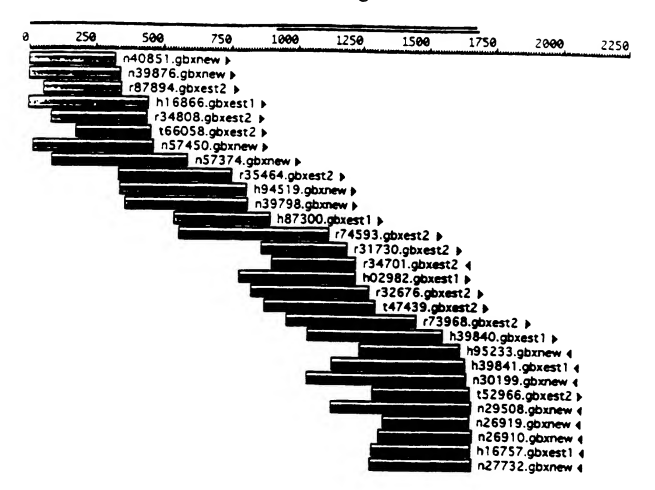
Figure 4 A.

Schematic depicting the overlap of ESTs bearing homology to the cDNA sequence encoding placental bikunin



Base pairs

Figure 4B



### Figure 4C

	L				50
Bikunin	GCGA	CCTCCGCSCS	TTGGGAGGTG	TAGCGCGGCT	CTGAACGCGT
440851		cctcccccc	TTGGGAGGTG	TAGEGEGEET	CTSAACGCGT
939876		cetecacaca	TTSSSASSITE	TAGCGCGGCT	CTSAACGCST
R87894					
H16866		cetecacaca	TTSGGAGGTG	TAGESES . CT	CTGAACGGGN
34808					
766058					
N57450				TAGCGCGGCT	CTGAACGCNA
N57374					
R35464					
H94519					
N39798					
H87300					
A74593					
R31730					
R34701					
H02982					
R32676	• • • • • • • • • • • • • • • • • • • •				
747439					
R73968	• • • • • • • • • • •				
H39840	• • • • • • • • • • • • • • • • • • • •	· · · · · · · · · · · ·			
H95233	• • • • • • • • • • • • • • • • • • • •	· · · · · · · · · · · · · · ·			
H39841	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • • • • •	• • • • • • • • • •	
N30199	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • • •		
752966	• • • • • • • • • • • • • • • • • • • •				
N29508	• • • • • • • • • • • • • • • • • • • •				
N26919	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • •	· · · · · · · · · · · · · · · ·		
N26910	• • • • • • • • • • • • • • • • • • • •	· · · • • • · · • · · · · · · · · · · ·	• • • • • • • • • •		
H16757	• • • • • • • • • • • • • • • • • • • •				
N27732	• • • • • • • • • • •	· · • • • • · · · · · ·	• • • • • • • • •		

	51-		•		100
Bikunin	GNA GGGCCS	TTSAGTSTCS	CAGGCGGCGA	SGGCGCGAGT	SAGSAGCAGA
N40851	NGAGNGGCCG	TTGAGTGTCG	CAGGGGGGGA	GGGCGCGAGT	SAGGAGCAGA
N39876	GCA.GGGGGG	TTGAGTGTCS	CAGGCGGCGA	GGGCGCGAGT	GAGGAGCAGA
387894		TTGAGTGTNG	NAGGEGGEGA	SGGCGCGAGT	GAGGAGCAGA
416066	ANGGGCCG	TTGAGTGTCG	CAGGEGGE . A	GGGCN . GAGT	GAGGAGCAGA
334808					GAGGAGCAGA
766058					
N57450	GAAGNGGCCG	TTGAGTGTCG	CAGGCGGGGA	SGGCGCGAGT	SAGGAGCAGA
N57374					AGA
R35464					
H94519					
N39798					
387300		• • • • • • • • • • • • • • • • • • • •			
R74593					
R31730					
R34701					
H02982					
232676				· · • • • • • • • • • • • • • • • • • •	
747439	• • • • • • • • • • • • • • • • • • • •				
a73968	• • • • • • • • • • • • • • • • • • • •	· · · · · · · · · · · · · · · ·			
H39040	• • • • • • • • • • •	• • • • • • • • •			
H95233	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • • •		
H39841	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	· · · · · · · · · · · ·		• • • • • • • • •
N30199	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •			
T52966	• • • • • • • • • •	• • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • •	
N29508	• • • • • • • • • • • •	• • • • • • • • • •			
N26919	· · · · · · · · · · · · · · · · · · ·	• • • • • • • • •			
N26910	• • • • • • • • • • •	• • • • • • • • •	• • • • • • • • • • •		
816757		• • • • • • • • • • •		· · · · · · · · · · · · · · · ·	
N27732				· · • • • • • • • •	

### 9/41

	101				150
Bikunin	CCCAGGCATC	GCGCGCGAG	AAGNO SSSS	STESSEACAC	TSAAGGTCCS
N40851	CCCAGGCATC	SCSCSCSAG	AAGNO . SSSC	GTCCCCACAC	TEAAGGTCCG
N39876	CCCAGGCATC	GCGCGCCGAG	AAGNO, SGGO	NTCCCCACAC	TSAAGGTCCS
287894	CCCAGGCATC	GCGCGCCSAG	AAGGCCGGGC	STCCCCACAC	TGAAGGTCCS
416866	CCCAGGCATC	SCGCGCCGAG	AAGNO . SEEC	GTCCCCACAC	TGAAGGTCCS
R34808	CCCAGGCATC	SCSCSCSAG	AAGNE . SSSC	STOCCCACAC	TGAAGGTCCG
T66058					
N57450	CCCAGGCATC	SCSTSCCSAG	AAGNO, SGGO	STECCEACAC	TGAAGGTCCS
NS7374	CCCAGGCATC	SCGCGCGGAG	AAGNO . SSSC	STEECCACAC	TGAAGGTCCG
R35464					
H94519					
N39798					
H87300					
274593					
R31730					
334701					
H02982					
R32676					
747439	· · · · · · · · · · · · · · · · · · ·				
973960					
H39840	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •			
H95233	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
939041	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • •		
N30199	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • •		
T52966	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •		
N29508	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	· · · · · · · · · · · · ·	
N26919	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •		
N26910	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
H16757	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •		• • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
N27732	• • • • • • • • • •	• • • • • • • • • •		• • • • • • • • • •	

	151				200
Bikunia	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	GGCGGACCCT	CCCCCACCCT
N40851	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	SSCSGACCCT	CCCGGAGCGT
N39876	GANAGGCGAC	ttccccccc	TTTGGCACCT	GGCGGACCCT	CCCGGAGCGT
387894	GAAAGGCGAC	TTTCGGGGGC	TTTGGCACCT	SSCSSACCCT	CCCGGAGCGT
H16866	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	SSCSSACS.T	CCCGGAGCN.
34808	SAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	SGCSGACCCT	CCCSSAGCST
766058				SGACCCT	CCCSGAGCGT
N57450	SAAAGGCSAC	TTCCGGGGGC	TTTGGCACCT	SGCSGACCCT	CCCSGAGCGT
457374	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	SSCGGACCCT	CCCSGAGCST
R35464					
H94519					
N39798	• • • • • • • • • • • • • • • • • • • •				
H87300	• • • • • • • • • • • • • • • • • • • •		· · · · · · · · · · · · · · ·		
374593					
R31730	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
R34701		• • • • • • • • • • •			
H02982		• • • • • • • • • • •			
R32676	· · · • • · · · · · · · · · · · · · · ·	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •		
747439		· · · · · · · · · · · · · · · ·	• • • • • • • • • • • • • • • • • • • •		
R73968		• • • • • • • • •			
H 3 9 8 4 0		• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • •
R 95233	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •		• • • • • • • • • • •	• • • • • • • • •
H39841	• • • • • • • • • •	• • • • • • • • • • •		• • • • • • • • •	
N30199	• • • • • • • • • • • •	• • • • • • • • • •		• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •
T52966	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •
N29508	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • •
N26919	• • • • • • • • • • • • • • • • • • • •	· · · · · · · · · · · · · · · ·	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •
N26913	• • • • • • • • • • • • • • • • • • • •			• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •
H16757	• • • • • • • • • • •	• • • • • • • • • •		· · · • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
N 2 7 7 32					

	201				250
Bikunin	CGGCACCTGA	ACGCGAGGCG	CTCCATTGCS	estacateta	. AGGGGGCTTC
N40851	CGGCACCTGA	ACGCGAGGCG	CTCCATTGCS	CSTGCGTNTG	. AGGGGCTTC
N39876	COCCACCTGA	ACGCGAGGCG	CTCCATTGCS	000000000000000000000000000000000000000	AGGGGCTTC
287894	CSGCACCTGA	ACGCGAGGCG	CTCCATTGCG	catacattta	AGGGGCTTC
416066	GGCACCTGA	ACGCGAGGCS	CTCCATTGCS	catacattts	AGGGGCTTC
34808	COGCACCTGA	ACGCGAGGCG	CTCCATTGCG	COTOCOTNIC	GAGGGGGTTC
766058	COSCACCISA	ACGCGAGGE.	DESTIGES	stacatata	NAGGGGGTTT
NS7450	COGCACCTGA	ACGCGAGGCG	CTCCATTGCG	catacattta	AGGGGCTTC
N57374	CSGCACCTSA	ACGCGAGGC.	STOCATION.	COTOCOTING	AGGGGCTTC
R35464					
H94519					
N39798					
H87300					
374593	· · · · · · · · · · · · · · · · · · ·				
<b>R31730</b>	• • • • • • • • • •				
R34701					
HC2982	• • • • • • • • • • • • • • • • • • • •				
R32676	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
T47439		• • • • • • • • • • • • • • • • • • • •			
R73968		· · · · · · · · · · · · · · ·			
439840		• • • • • • • • • •			
H95233	• • • • • • • • • • • • • • • • • • • •				
H39841	• • • • • • • • • • • • • • • • • • • •				
N30199	• • • • • • • • • • • • • • • • • • • •	· · · · · · · · · · · · · · ·			
T52966	• • • • • • • • • •		· · · · · · · · · · · ·		
N2 9508	• • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •		
N2 6919	• • • • • • • • • • • • • • • • • • • •	· · · · · · · · · · · · ·			• • • • • • • • • • •
N2 6910					
H16757	• • • • • • • • • • •				
N27732		· · · · · · · · · · · ·		<b></b>	

•	_	•			
	251				300
Bixunin	CCGCACCT G	ATCGCGAGAC	CCCAACGGCT	satss care	se to cocs
N40851	CCCCACCT.G	ATCCCSAGAC	CCCAACSGCT	SSTSS.CSTC	scets.cses
N39876	CCSCACCT.G	ATCCCGAGAC	CCCAACGGCT	sates.cate	scata.cscs
387894	CCGCACCT.S	ATCGCGAGAC	CCCAACGGCT	SGING.CGIC	SC.TN.CGCS
H16866	CCGCACCT.G	ATCCCGAGAC	CCCAACGGCT	SGING.COTC	sc.tsscscs
334808	COSCACCT.S	ATCGCGAGAC	CCCAACGGCT	SCTSSSCSTE	sc.ts.cscs
766058	CCSCACCT.S	ATCGCGAGAC	CCCAACGGCT	30150.0010	sc.ts.cscs
N57450	COSCACCT.S	ATCCCGAGAC	CCCAACGGCT	\$6756.0570	scets.eses
N57374	CCGGAACTTG	ATCGCGAGAC	CCCAACGGCT	SSTSS.CSTC	GC.TG.CGCS
335464					
H94519					
N39798					
H87300					
R74593					
R31730					
R34701					
H02982					
R32676					
T47439				· · · · · · · · · · · · · · · · · · ·	
R73968					
H39840					
H95233					
H39841				• • • • • • • • •	
N30199					
T52966					
N29500					
N26919					
N26910					
H16757					
N27732	· · · · · · · · · · · · · · · · · · ·				

	301				350
31kunin	דכ זכבבכדב	AGCT SSCCA	TOGCGCANT	stice seec	T SAGGE GS
940851	TC.TCSGCTG	AGCT.SGNCA	79 <b>7</b> 00		
939876	TC.TCGGCTG	AGCT.GGCCA	TOGCOCACT.	G. TGCGGNGC	T. SAGGE.S
387894	TC.TCGGCTG	AGCTTGGCCA	TOGCOCANT.	STTNE.SGGC	T. NAGGO. SG
H16866	TTCTCSGCTG	AGCT. SGCCA	TOGCOCANT.	STTSC. GNGC	T.GAGGC.GG
R34808	TOTTCSSCTS	AGCTGGGCCA	TGGCGCANTT	STTGC.SGGC	T.GAGGE.SS
166058	TC.TCSGCTG	AGCT.SGCCA	TOGCOCANT.	STTSC. SNGC	T.GAGGE.GS
N57450	TC.TCGGCTG	AGCT, GGCCA	TOOCGCANT.	GSTSC.SGSC	TTGAGGE.GG
N57374	TOOTOSSOTS	AGCT. GGCCA	TGGCGCANT.	SSTGCCGNGC	T. SAGGCCGG
R35464					
R94519		• • • • • • • • • • •			
N39798					
H87300					
274593					
R31730					
R34701	• • • • • • • • • • •		· · · · · · · · · · · ·		
H02982	• • • • • • • • • • • • • • • • • • • •				
R32676	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •		• • • • • • • • • •	
T47439			· · · · · · · · · · · · · · ·		
373968	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • • • • • • • • • • • •		
939640	• • • • • • • • • •		• • • • • • • • • •	• • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
H95233	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •
H39841	• • • • • • • • • •			• • • • • • • • • •	• • • • • • • • •
NJ0199		· · · · · · · · · · · · · · · · · · ·	• • • • • • • • •	• • • • • • • • • • •	
T52966	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •
N29508	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • •
N26919	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •	• • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
N26910	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •	• • • • • • • • • •	• • • • • • • • • •
H16757		• • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
NZ / / 32		<b></b>	<b>.</b>		

•	351				400
Bikunin	AC GG CG	TTTCTCS	00 7507566	A TOSCT SC	: ::::::::
R87894	ACS.				
H16866	AC CGNCGT	TTTTCTTCS.	contactage	ATTOSOTTEC	TTECTNTCTS
334808	ACCCCONCO.	. ITTTTTCSN	COTTSCTGGG	ATTCS.TTS.	TENCTOTOTA
T66058	CSGNCG.	. TTTTCTCS.	co.toctood	Autocotuse	t.cototot.
N57450	ANN. NGCCC.	: : : : : : : : : : : : : : : : : : :	00.7007666	ALTOSET . SC	T. SETETET.
N57374	AGSSCCSG	:ttctca.	cottoctoss	ALTOSCT.SC	t.corcrets
R35464	gtcg.	TTTCTCG.	corssoress	A.TOSCT.SC	T.CCTCTCT.
H94519	SCNGCGCG.	ITNNTCG.		ALTOGOT.SC	
N39798			<b>ctsg</b>	ANTOGOT, SC	T.CCTCTCT.
H87300		· · · · · · · · · · · · · · · · · · ·			
R74593	• • • • • • • • • • • • • • • • • • • •				
831730		· · · • • • · · · · · · · · · · · · · ·			
R34701					
HC2982					
<b>32676</b>	• • • • • • • • • • • • • • • • • • • •	· • • • • • • • • • • • • • • • • • • •			
747439					
R73968					
H39840					
H95233	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	· · · · · · · · · · · ·	· · · · · · · · · · · · ·	• • • • • • • • • •
H39841		• • • • • • • • • • • • • • • • • • • •			· · · · · · · · · · · · · · · ·
ИЭО199				• • • • • • • • • • • •	
752966				• • • • • • • • • • • • • • • • • • • •	
N29508			· · · · • • • • • • • •		• • • • • • • • •
N2 6919		• • • • • • • • • • •		• • • • • • • • • • • • • • • • • • • •	· · · · · · · · · · · · · · · ·
N26910		· · · · · · · · · · · · ·		• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •
H16757				• • • • • • • • • • • • • • • • • • • •	
N27732					

	401	a
Bikunin	GGGG TCCTG G CGGCCGA CCGA GAACG CA GCA TCC ACGACTT C	Ţ
H16066	GOOGTTCCTG GG.COGCCGA CCGA.GAACG CA.GCA TOC AAGAATTTT	
R34808	GGGGTTC.TG GGGNGGCCCA NCGA.GAACG CAAGCA.TTC ACGA.TTT	•
T66058	GOGG. TOOTS G COGGCCGA CCGA.GAACG CA.GCA.TCC ACGANTT.C	+
N57450	GGGG.TCCTG GCGGCCGA CCGA.GAACG CA.GCA.TCC ACGACTT.C	
N57374	GGGG. TEETS GCGGCCGA NCGAAGAANG CA.GCAATCC ANGAATTNC	
R35464	GGGG. TOOTG G.CCGGGCCGA CCGA.GAACG CA.GCA.TCC ACGACTT.C	
H94519	GGGG. TOGNG GCGGCCGA CCGA.GAACG CA.GCA.TCC ACGACTT.C	
H39798	GGGG. TOOTS G CGGCCGA CCGA.GAACG CA.GCA. TCC ACGACTT. C	
H87300		
R74593	***************************************	
<b>331730</b>		
R34701		
H02982		
R32676		
T47439		
R73968		
H39040		
H95233		
H39841		
N30199		
T52966		
N29508		
N26919		
N2 6910		
H16757 N27732		
74//12		

	451				500
atnuxte	GCCTGGTGT	CGAAGGT SS	TGGGGAGATG	00000 0000	CATGCCTA G
H16866	GCC				
T66058	TECTEGTETT	CGAAGG			
N57450	SCCTSSTST.	CGAAGGT.GG	TOGGCAG		
<b>357374</b>	SCCTSSTGTT	CCAAAGTTGG	TOGGCANATT	cosssscent	CATGNOTAAG
335464	SCCTSGTGT.	CGAAGGT.GG	TOGGCAGATT	00000.0000	CATGCCTA.G
H94519	SCCTSSTGT.	CGAAGGT.GG	TOGGCAGATG	ccccc.cctc	CATGCCTA.G
<b>39798</b>	SCCTSSTGT.	CGAAGGT.GG	TGGGCAGATG	ccasa.cctc	CATGCCTA.G
887300					
R74593					
R31730					
834701					
H02982	· · · · · · · · · · · · · · · · · · ·				
332676	• • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
747439	• • • • • • • • •				
273968				· · · · · · · · · · · · ·	
H39840	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •		· · · · · · · · · · · · · · · ·	
195233	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •			
H39041	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •			
N30199	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • •		
752966	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
N29508	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
N26919	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • •	• • • • • • • • • •		
N26910	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
H16757	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •			
N27732			. <b>. .</b>		

	9010 10	(COH C	• •		
	501				550
Bikunin	c test set	ACAATGTCAC	TSACGGATCS	TSCCAGCTST	TESTSE ATS
N57374	ctt <del>cc</del> ttcct		TTAANGATTI	TISCANCIST	TIGIGINATI
R35464	G.TSST.SGT	ACANTGICAC	TGACGGATCC	TGCCAGCTST	TISTGT.ATG
H94519	G . TOST . SST	ACANTGTEAC	TSACSGATCC	TSCCAGCTST	TTSTST.ATS
N39798	c.tsst.sst	ACAATGTCAC	TSACGGATCS	TSCCAGCTST	TTGTGT.ATG
487300		• • • • • • • • • • • • • • • • • • • •			
R74593					
331730					
334701					
H02982					
R32676					
T47439					
R73968					
H39840		• • • • • • • • • • • • • • • • • • • •			
495233					
HJ9841					
N30199					
T52966					
N29508					
N2 6919					
N2 6910	· · · · · · · · · · · · · · · · · · ·				
H16757	• • • • • • • • • • • • • • • • • • • •				
N27732	• • • • • • • • • • • • • • • • • • • •				
	551				600
Bikunin	CCCCCTCTCA			CCTGACCAAG	
N57374	GGGGCTNTTA	AACGGAAANA	.CAATAATNA	CCTGACCAAA	GAAGNAAT
R35464	CCCCCTCTCA		GCAATAATTA	CCTGACCAAG	CA.GCAGTGC
H94519	GGGGCTGTGA	COGAAACA	GCAATAATTA	CCTGACCAAG	SA.GGAGTGC
N39798	CCCCCTCTCA	CGGAAACA	GCAATAATTA	CCTGACCAAG	GA . GGAGTGC
H87300	GATTCSSCAC	AGGGGAAACA	GCAATAATTA	CCTGACCAAG	GA . GGAGTNC
R74593	• • • • • • • • • • • •	• • • • • • • • •	SCAATAATTA	CCTGACCAAG	GA.GGAGTGC
R31730	• • • • • • • • • • • • • • • • • • • •	· • • • • • • • • • • • • • • • • • • •			• • • • • • • • • • • • • • • • • • • •
R34701	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •		• • • • • • • • • • •
H02982	· • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
R32676	• • • • • • • • • • •	• • • • • • • • •			
747439	· · · · · · · · · · · · · · · · · · ·				
R73968	•••••	• • • • • • • • • • • • • • • • • • • •			
H39040	• • • • • • • • • • • • • • • • • • • •				
H95233	•••••				
H39641	• • • • • • • • • • • • • • • • • • • •				
N30199	• • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
T52966					
N29508	•••••	• • • • • • • • • • •		· · · · · · · · · · · · ·	
	••••••	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	
N26919					
N26919 N26910		•••••		• • • • • • • • • • • • • • • • • • • •	
				• • • • • • • • • • • • • • • • • • • •	

* * '	yara ic	(Con c	,		
	6Q1				653
grknuru	CTCAAGAAAT	STSCCACTST	CACAGAGAAT	SCCACGGGTG	ACCTOUCCAC
R35464	CTCAAGAAAT	STECCACTET	CACAGAGAAT	SCCACGGGTG	ACCESCEAG
H94519	CTCAAGAAAT	STGCCACTST	CACAGAGAAT	SCCACGGGTG	ACCTOGCCAC
N39798	CTCAAGAAAT	GTGCCACTGT	CACAGAGAAT	SCCACGGGTS	ACCTGGCCAC
d87300	CTCAAGAAAT	STNCCACTGT	CACAGAGAAT	GCCACGGGTS	ACCTOSCOAC
R14593	CTCAAGAAAT	STSCCACTST	TACAGAGAAT	SCCACGGGTS	ACCTGGCCAC
231730					
R34701					
HQ2982					
R32676					
147439					
373968					
839840					
H95233					
H39641					
N30199					
752966					
N29508					
N26919					
N26910					
H16757					
N27732					
	651				700
Bikunin	CAGCAGGAAT	GCAGCGGATT	CCTCTGTCCC	AAGTGCTCCC	AGAAGGCAGG
R35464	CAGCAGGAAT	GCAGCGGATT	CCTCTGTCCC	AAGTGCTCCC	AGAAGGCAGG
H94519	CAGCAGGAAT	GCAGCGGATT	CCTCTGTCCC	AAGTGCTCCC	AGAAGGCAGG
N39798	CAGCAGGAAT	SCAGCGGATT	CCTCTGTCCC	AAGTGCTCCC	AGAAGGCAGG
H87300	CAGCAGGAAT	GCAGCGGATT	CETETETECE	AAGTGCTCCC	AGAAGGCAGG
R74593	CAGCAGGAAT	SCAGCSGATT	CCTCTGTCCC	AAGT.CTCCC	AGAAGGCAGG
R31730		· · · · · · · · · · · ·			
R34701	• • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
H02982	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • •	
932676	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	
T47439		• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	· • • • • • • • • • • • • • • • • • • •	· · · · · · · · · · · · · · · · · · ·
R73968	• • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • •
R39840	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			• • • • • • • • • • • • • • • • • • • •
H95233					
a39441	•••••				
N30199	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •			
752966	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
N2 9508	• • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
N26919					
N26910		• • • • • • • • • • • • • • • • • • • •			
H16757					•••••

	701	•			
91kunin	• -	ACCACTOCA			750
R35464		ACCACTICAC			AAGAATACTG
94519					AAGAATAATT
N39798		; ACCACTCCAC ; ACCACTCCAC			AAGAATACTG
487300		. ACCACTCCAC		CAACTATG	AAGAATACTG
R14593		ACCACTOCAC			AAGAATACTS
R3:130					AAGAATACTG
R34701					********
H02982					
332676					
747439	• • • • • • • • • • • • • • • • • • • •				********
R73968	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • • • • • • • • • • • •
H39840					• • • • • • • • • • • • • • • • • • • •
H95233	• • • • • • • • • • • • • • • • • • • •			• • • • • • • • • •	• • • • • • • • • • •
H39841	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
N30199	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • • •	
752966		• • • • • • • • • • • • • • • • • • • •	- · · · · · · · · · · · ·	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •
N29508	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • •	• • • • • • • • •	• • • • • • • • •
N2 6919	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • •		• • • • • • • • •
N26919	• • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •		• • • • • • • • • • •
H16757	*********	• • • • • • • • • • •		• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
N27732	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
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	751				
Bikunin		CCCACT CAC	TGGGCC TTG		100
R35464	GCACCGNCAA		roote iis	CCGIG CAT	CCTT CCCAC
H94519					
N39798	JENE COCCION				
	CACCCCCAA	CCCACT CAC		C.GTG.CAT.	
HB/300	. CACCGCCAA	CGCAGT.CAC	TSGSGCCTTS	C.GTGGAAT.	CCTTTCCCAC
H87300 R74593	.CACCGCCAA	CGCAGT.CAC CGCAGTNCAC	TSGGGCCTTG TSGGCC.TTG	C.GTGGAAT.	CCTTTCCCAC
R74593	.CACCGCCAA	CGCAGT.CAC CGCAGT.CAC	TSGGGCCTTG TSGGCC.TTG TSGGGC.TTG	C.GTGGAAT. C.GTGGCATN CCGTG.CAT.	CCTTTCCCAC CCTT.CCCAC
R74593 R31730	CACCGCCAA CACCGCCAA	CGCAGT.CAC CGCAGTNCAC CGCAGT.CAC	TSGGGCCTTG TSGGCC.TTG TGGGCC.TTG	C.GTGGAAT. C.GTGGCATN CCGTG.CAT.	CCTTTCCCAC CCTT.CCCAC
R74593 R31730 R34701	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGTNCAC CGCAGT.CAC	TSGGGCCTTG TSGGCC.TTG TSGGCC.TTG	C.GTGGAAT, C.GTGGCATN CCGTG.CAT.	CCTTTCCCAC CCTT.CCCAC
R74593 R31730 R34701 H02982	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGT.CAC	TSGGGCCTTG TSGGGCC.TTG TSGGGCC.TTG	C.GTGGAAT.	CCTTTCCCAC CCTT.CCCAC
R74593 R31730 R34701 H02982 R32676	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGTNCAC CGCAGT.CAC	TGGGGCC.TTG TGGGGCC.TTG	C.GTGGAAT.	CCTTTCCCAC CCTT.CCCAC
R74593 R31730 R34701 H02982	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGTNCAC CGCAGT.CAC	TGGGGCC.TTG TGGGGCC.TTG	C.GTGGCATN (	CCTTTCCCAC CCTT.CCCAC CCTT.CCCAC
R74593 R31730 R34701 H02902 R32676 T47439 R73960	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGTNCAC CGCAGT.CAC	TSGGGCCTTG TSGGGCC.TTG TSGGSCC.TTG	C.GTGGAAT.	CCTTTCCCAC CCTT.CCCAC
R74593 R31730 R34701 H02902 R32676 T47439 R73960 H39840	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGTNCAC CGCAGT.CAC	TSGGGCCTTG TSGGGCC.TTG TSGGSCC.TTG	C.GTGGAT.	CCTTTCCCAC CCTT.CCCAC
R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGTNCAC CGCAGT.CAC	TSGGGCCTTG TSGGGCC.TTG TSGGSCC.TTG	C.GTGGAT.	CCTTTCCCAC CCTT.CCCAC
R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233 H39841	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGT.CAC	TSGGGCCTTG TGGGCC.TTG TGGGCC.TTG	C.GTGGAT.	CCTTTCCCAC CCTT.CCCAC
R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGT.CAC	TSGGGCCTTG TSGGGCC.TTG TSGGSCC.TTG	C.GTGGCATN CCGTG.CAT.	CCTTTCCCAC CCTT.CCCAC
R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 K95233 H39841 N30199	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGT.CAC	TSGGGCCTTG TGGGCC.TTG	C.GTGGAAT.	CCTTTCCAC CCTT.CCCAC
R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233 H39841 N30199 T52966	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGTNCAC CGCAGT.CAC	TSGGGCCTTG TGGGCC.TTG TGGGCC.TTG	C.GTGGAT.	CCTTTCCAC CCTT.CCCAC
R74593 R31730 R34701 H02902 R32676 T47439 R73960 H39840 H95233 H39041 N30199 T52966 N29500 N26919	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGTNCAC	TSGGGCCTTG TGGGCC.TTG TGGGCC.TTG	C.GTGGAT.	CCTTTCCAC CCTT.CCCAC
R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233 H39841 N30199 T52966 N29508 N26919	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGTNCAC CGCAGT.CAC	TSGGGCCTTG TGGGCC.TTG TGGGCC.TTG	C.GTGGAT.	CCTTTCCAC CCTT.CCCAC
R74593 R31730 R34701 H02902 R32676 T47439 R73960 H39840 H95233 H39041 N30199 T52966 N29500 N26919	.CACCGCCAA .CACCGCCAA	CGCAGT.CAC CGCAGT.CAC	TSGGGCCTTG TGGGCC.TTG TGGGCC.TTG	C.GTGGAT.	CCTTTCCAC CCTT.CCCAC

		(COH C			
	801				850
Bikunin	GCTGGTACTT	T GACGTSSA	GA GGAACTO	CTS CAATAA	CTICATCTAT
H94519	GCTGGTACTT	TIGNOST			
N39798	GCTGGNAATT	THEACETTEA	GAAGGAAC		
H87300	GCTNGTACTT	T. SACSTSSA	GA. GGAACTI	STSSCAATAA	CTTCATCTAT
374593	GCTGGTACTT	T.GACGTGGA	SA.GGAACTC	CTG. CAATAA	CTTCATCTAT
R31730					
R34701					
H02982			GA. SGAACTC	CTG.CAATAA	CTTCATCTAT
232676				3	ATTCSGAA
147439					
R73968					
H39840					
H95233					• • • • • • • •
H39841					
N30199				• • • • • • • • • • • • • • • • • • • •	
752966				· · · · · · · · · · · · · · · · · · ·	• • • • • • • • • •
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N26919					· · · · · · · · · · · · · · ·
N26910	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • •		
H16757			• • • • • • • • • •	• • • • • • • • • • •	• • • • • • • • • •
N27732	• • • • • • • • • •				•
	851				
	937				900
Bikunin	GGAGGCT GC			TACCGCTC T	GAGGAGGCCT
H87300	GGAGGET GE GGAGGETTGE	CSSSSCAATN	AAGAACAGNT	TACCGCTCTT	GAGGAGGCCT TAGGAGGCCT
H87300 R74593	GGAGGET GE GGAGGETTGE		AAGAACAGNT AAGAACAG.C	TACCGCTCTT TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730	GGAGGET GE GGAGGETTGE GGAGGET.GE	CSSSSCAATN	AAGAACAGNT AAGAACAG.C	TACCGCTCTT	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701	GGAGGCT GC GGAGGCTTGC GGAGGCT.GC	CSGSGCAATN CSGGGCAAT.	AAGAACAGNT AAGAACAG.C G.C	TACCGCTCTT TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701 H02982	GGAGGCT GC GGAGGCTTGC GGAGGCT.GC	CGGGGCAATN CGGGGCAAT. CGGGGG.AAT.	AAGAACAGNT AAGAACAG.C G.C AAGAACA.NC	TACCGCTCIT TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCTGAGGAGGCCT
H87300 R74593 R31730 R34701 H02982 R32676	GGAGGCT GC GGAGGCT.GC	CGGGGCAATN CGGGGCAAT. CGGGG, AAT. CGGGGCAAT.	AAGAACAGNT AAGAACAG.C G.C AAGAACA.NC AAGAACAG.C	TACCGCTCTT TACCGCTC.T TACCGCTC.T TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701 H02982 R32676 T47439	GGAGGCT GC GGAGGCT GC	CGGGGCAATN CGGGGCAAT. CGGGGG.AAT.	AAGAACAGNT AAGAACAG.C G.C AAGAACA.NC AAGAACAG.C	TACCGCTCTT TACCGCTC.T TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701 H02982 R32676 T47439 R73968	GGAGGCT GC GGAGGCT.GC	CGGGGCAATN CGGGGCAAT. CGGGGCAAT.	AAGAACAG.CG.CAAGAACA.NC	TACCGCTCTT TACCGCTC.T TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840	GGAGGCT GC GGAGGCTTGC GGAGGCT.GC	CGGGGCAATN CGGGGCAAT. CGGGGCAAT.	AAGAACAG.CG.CAAGAACA.NC	TACCGCTC.T TACCGCTC.T TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233	GGAGGCT GC GGAGGCT.GC GGAGGCT.GC GGAGGCT.GC	CGGGGCAATN CGGGGGAAT. CGGGGGAAT.	AAGAACAGNT AAGAACAG.C G.C AAGAACAG.C AAGAACAG.C	TACCGCTC:T TACCGCTC:T TACCGCTC:T TACCGCTC:T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233 H39841	GGAGGCT GC GGAGGCT.GC GGNGGCT.GC CGAGGAGC	CGGGGCAATN CGGGGCAAT. CGGGGG.AAT. CGGGGGCAAT.	AAGAACAGNT AAGAACAG.C G.C AAGAACAG.C AAGAACAG.C	TACCGCTC.T TACCGCTC.T TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233 H39841 N30199	GGAGGCT GC GGAGGCT GC GGNGGCT GC GGNGGCT GC	CGGGGCAATN CGGGGCAAT. CGGGGCAAT.	AAGAACAG.CG.CAAGAACA.NC	TACCGCTC:T TACCGCTC:T TACCGCTC:T TACCGCTC:T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233 H39841 N30199 T52966	GGAGGCT GC GGAGGCT GC GGAGGCT GC GGNGGCT GC	CGGGGCAATN CGGGGCAAT. CGGGGCAAT.	AAGAACAGNT AAGAACAG.C G.C AAGAACAG.C AAGAACAG.C	TACCGCTCTT TACCGCTC.T TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233 H39841 N30199 T52966 N29508	GGAGGCT GC GGAGGCT GC GGAGGCT GC GGAGGCT GC	CGGGGCAATN CGGGGCAAT. CGGGGCAAT.	AAGAACAG.CG.CAAGAACAG.C	TACCGCTC.T TACCGCTC.T TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233 H39841 N30199 T52966 N29508 N26919	GGAGGCT GC GGAGGCT GC GGAGGCT GC GGAGGCT GC GGAGGCT GC	CGGGGCAATN CGGGGG.AAT. CGGGGG.AAT.	AAGAACAG.CG.CAAGAACAG.C	TACCGCTC.T TACCGCTC.T TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT
H87300 R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233 H39841 N30199 T52966 N29508 N26919 N26910	GGAGGCT GC GGAGGCT GC GGAGGCT GC GGAGGCT GC GGAGGCT GC	CGGGGCAATN CGGGGG.AAT. CGGGGG.AAT.	AAGAACAGNT AAGAACAG.C G.C AAGAACAG.C AAGAACAG.C	TACCGCTC.T TACCGCTC.T TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCTBGGCCT
H87300 R74593 R31730 R34701 H02982 R32676 T47439 R73968 H39840 H95233 H39841 N30199 T52966 N29508 N26919	GGAGGCT GC GGAGGCT.GC GGAGGCT.GC GGAGGCT.GC CGAGGAGC	CGGGGCAATN CGGGGG.AAT. CGGGGG.AAT.	AAGAACAGNT AAGAACAG.C G.C AAGAACAG.C AAGAACAG.C	TACCGCTCTT TACCGCTC.T TACCGCTC.T TACCGCTC.T	GAGGAGGCCT TAGGAGGCCT GAGGAGGCCT GAGGAGGCCT GAGGAGGCCT

ri	gure	4 C	(Con'	t)		
	901					950
Bikunin	GCA	TGCTO	cscrscrrc	c sc		CA SCAGGA
H87300		Ī				The School
R74593	. SCA.	TGCTC	cactactic	c		CA . GCAGGA
831730	.SCA.	TSCTC	cactacttc	c sc		
R34701				c		- CAAGCAGGA
H02982	. 305.	tacto	coctoctto			
R32676	SCA.	tsete	cactactto:			
T47439	TSCAG	TGCTC	CGCTGCTTC	: ac		
273968		<b>.</b> .			·	
H39840						
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T52966						
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H16757	· · · · · ·					
N27732		• • • •	• • • • • • • • • • •			
866	951					1000
Bikunin R74593	CAA TO	TTCC :	000000011	GGCTCAAAGG	TESTESTIC	TGG CGGGGC
R31730				GGCTCAAAGG		೨ <b>೦೦೦೦೦೦೦</b>
R34701		STCC (	CTGCCCCTT	GGCTCAAAGG	TGGTGGTTC.	TGG.CGGGGC
H02982	AAANTC			GGCTCAAAGG	TGGTGGTTCC	TGG.CGGGGC
R32676	GAA. TCC				TGGTGGTTC.	TGG.CGGGGC
T47439			CTGCCCCTT		TGGTGGTTC.	TGG.CGGGGC
R73968	SAA. TCC		CTGCCCCTT	GGCTCAAAGG	TGGTGGTTC.	TGG.CGGGC
H39840	• • • • • • •		• • • • • • • • • • •		• • • • • • • • • • • • • • • • • • • •	с66666
H95233		• • •	• • • • • • • • •	• • • • • • • • • •		
H39841		• • • •	• • • • • • • • • •	• • • • • • • • • • •		
N30199		• • • •		• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	
752966				• • • • • • • • •	• • • • • • • • • •	
NZ 9508		•••	• • • • • • • • •		• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •
N26919		• • • ·				
N26910				· · · · · · · · · · · · ·	• • • • • • • • • •	
H16757					• • • • • • • • • • • • • • • • • • • •	
N27732			• • • • • • • • • •			

	y 10	(COH	<b>C</b> )		
	1201				: 050
Bikunin	TGTT CGTGA	TGGTGTTGA	T 00 T 0TTC	TSGS AGCC	T DO ATSOTO
R74593	TGTTTCGTGA	TGGTGTTSA	: cott:tc	TOSSSAGEN	T CC.ATGGTCT
R31730	TGTT.CGTGA	TGGTGTTGA	T colilette	: TOGGGAGCC	CC.ATGGTC.
234701	TSTT.CGTGA	TEGTETTEA	<b>:</b> :::::::::::::::::::::::::::::::::::	cood. Adde:	CCCATGGTCC
HC2982	TGTT.CGTGA	TGGTGTTGA	: 00.7.0 <b>77</b> 06	TGGG . AGCC:	CC.ATCGTN.
332676	TGTT . CGTGA	TGGTGTTGA	1 00.1.01100	TOGG . AGCC:	
747439	TOTT. COTOX	TGGTGTTGAT	r colticated	TGGG . AGCC:	CO.ATOGTO.
373968	TOTT.CGTGA	TESTSTTEAT	7 00.7.07 <b>700</b>	TOGG . AGCCT	CC.ATCSTC.
839840	• • • • • • • • • • •				
H95233	• • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
H39841		· · · · · · · · · · · ·			
N30199	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
T52966		• • • • • • • • • • • • • • • • • • • •			
N2 9508		• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •		
N26919	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
N26910	• • • • • • • • • • •	• • • • • • • • •			
416757	• • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
N27732	• • • • • • • • • • •	• • • • • • • • • •			
	. 051				
Bikunin	1051				1100
R74593	TACC TOAT	CCSCGTGGCA	COGAGG AAC	C AGG AGCG	TGCCCTGCGC
R31730	TAC. TGATT				TGCCCTGCGG
R31730			CCCACCCAAC		TGCCCTGCGC
HO2982			CGGAGG . AAC		TGCCCTGCGC
R32676			CGGAGG. AAC		TGCCCTGCGN
T47439			CGGAGG . AAC		TGCCCTGCGC
R73968			CGGAGG. AAC		TGCCCTGCGC
H39840			CGGAGG . AAC		TGCCCTGCGC
H95233			GGG.AAC	C. AGG. AGCG	TGCCCTGCGC
H39841		• • • • • • • • • • • • • • • • • • • •			• • • • • • • • •
N30199		• • • • • • • • •			
T52966		• • • • • • • • •	GAGGAACC		TECECTGEGE
N2 9508		• • • • • • • • • •	••••		• • • • • • • • •
N26919		• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
N26910		• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •		
H16757		• • • • • • • • •		· · · · · · · · · · · · · · · ·	• • • • • • • • • • • • • • • • • • • •
N27732		• • • • • • • • • • • • • • • • • • • •			• • • • • • • • • •
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	<b>3</b> 10	(COM )	- /		
	1101				1150
Bikunin	ACCS TOT S	SAGCTCUSS.	A JATSACAAGS	AGCAGCTSS	TURAGARO
R74593	ANCG.TCT.S	SASCTTESS	A SATSACAASS	SNT	
831730	ACCG.TCTGG	SAGCTCCSS	A GATGACAAGG	SAGGAGGTGG	STSAAGAAC.
334701	ACCS.TCT.G	CAGCTCCCC	A GATGACAAGG	. AGCAGCTGG	.TSAAGAAC.
402982			N SATSACAAGG	. AGCAGCTGG	TSAAGAAC.
732676	ACCC.TCTSS	SAGCTCCGG	A SATSACAASS	GAGCASCISS	TSAAGAAC.
747439	ACCC.TCT.C	SAGCTCCGG	N GATGACAAGG	AGCAGCTGG	. TGAAGAAC .
R:3968	ACCG.TCT.G	SAGCTCCSGA	N SATSACAAGS	AGCAGCTGG	TGAAGAAC.
439840	ACCESTET . S	SAGCTCCSGA	SATGACAAGG	AGCAGCTGG	TGAAGAAC.
495233					
H39841					
N30199	ACCG.TCT.G	GAGCTCCSGA	GATNACAANG	AGCAGCTON	TUAAGAACC
752966					
N29508					
N26919			• • • • • • • • • •		
N26910					
H16757					
N27732					
•	1151				1200
Bikunin			ccctst ccc		
R31730			NCCTCTTCCC		
R34701			ccctgt.cgc		
H02982			NCCTGTTCGN		
R32676 T47439			CCCTGTTCGC		
R73968			CCCTGT.CGC		
H39840			CCCTGT.CGC		
H95233			CCCTGT.CGC		ET.NGGGAA.
H39841			• • • • • • • • • • • • • • • • • • • •		
N30199			CCCTGT.CGC		
752966	ACATATGT.C				
N29508					
N26919			CCCTNT.CGC (		T.SGG.AA.
N26919 N26910					
H16757					• • • • • • • • • • • • • • • • • • • •
N27732				· · · · · · · · · · · · · · · · · · ·	
74 1 1 32					

Fi	gure 4C	(Con't	.)		
	1201				.250
Bikunin	GGGAGGGG	AGACTAT S	TOT GA GCT	AA	A TAGA SG
831730	GGGAGGGGG	A			,30
R34701	GGGAGGGG.	AGACTAT.S.	דבד בא. בכד	TTTTTTAA	A *A
H02982	GGGGAGGG	AGATTAT S.			
932676	SCOGAGGGG				
747439	SSSASSSS.	AGACTAT.S.			ATTAGGAGGG
373968		AGACTAT.S.			A.TAGAGG
139840		AGACTAT.S.			A.TAGAGG
195233					A.TAGAGG
H39841			TGT GAACCT		
N30199				•	A.TAGA,.GG
T52966				TITTITAA	
N29508					
N26919				TTTTTTAA	
N26919					
H16757					
N27732				· · · · · · · · · · · · · · · · · · ·	
427.32				• • • • • • • • • •	
	1251				
Bikunin	GATTGACTC				: 300
R32676			GT GATC A		GAGGTCTGTT
T47439			GTTGATCCAT	TTAGGGGGGNT	
773968			ST.SATC.A.	TTAGGGCT	
H39840			GT.SATC.A.		
H95233			GT.GATC.A.		
H39641 N30199			GT.GATC.A.		
752966				TTAGGGCT	GAGGTCTGTT
N29508 N26919				TTAGGGCT	
N2 6910			• • • • • • • •		
H16757					
N27732				· · · · · · · · · · · · · · · · · · ·	
<b>5</b> : <b>4</b>	1301				1350
Bikunin	TCTCTGGGAG G		TGCTTCC TG	S TO TOGGA	SGGATGGG
T47439					
R73968	TCTCTGGGAG G				
H39840	TCTCTGGGAG G				
H95233	NCTCTGGGAG N				
H39841	TCNCTGGGAG G				
N30199	TCTCTGGGAG G				
752966					
N29508	TETETGGGAG G				
N26919		• • • • • • • • •			
N26910	• • • • • • • • • • •				
H16757					
N27732	÷	· • • • · · • · ·		GGTCCTGNCA	AGGNATGGGG

```
Figure 4C (Con't)
          1351
                                                          1400
 Bikunin TTTG CTTTG G AAATCCTC T AGGAGGCT CCTCCT CGC ATGG CC TG
  RR3968 TITG.CITTG GGAAATCOTG TINGGAGGCT COTCOTTOGC ATGGGCCTTG
  H39840 TTTG.CTTTG GAGAATCCTC T ANGAGGET COTCCT.CCC ATGG.CC.TG
  H95233 TTTG.CTTTG G.AAATCCTC T.AGGAGGCT CCTCCT.CGC ATGG.CC.TG
  H39841 TTTG.CTTTG G.AAANCONG T.AGGAGGET COTCCT.CGC ATGG.CC.TG
  N30199 TTTG.CTTTG G.AAATGGTG T.AGGAGGGT GGTGGTTGGG ATGG.CG.TG
 T52966 TTTG.CTTTG G.AAATCCTC T.AGGAGGCT CCTCCT.CGC ATGG.CC.TG
 N29508 TITG.CTTTG G.AAATCCTG T.AGGAGGGT GCTCCT.CGC ATGG.CC.TG
 N26919 ..... ATGG.CC.TG
 N26910 .....CTTTT GNAAATCGTG T.AGGAGGGT CCTCCT.CGC ATGG.CC.TG
 H16757 TTTGCCTTTG G.AAANCCTC T.AGGAGGCT COTCCT.CGC ATGG.CC.TG
 N27732 TTTG.CTTTG G.AAATCCTC TTAGGAGGCT CCTCCT.CGC ATGG.CC.TG
         1401
                                                         1450
Bikunin CAGT CT GG CAGCAG CCC CGAGTTGTTT CC TCGCTG ATC GATTTC
 R73968 CAGT.CTNGG CAGCANCCCC CGAGTTTTTT TCCTTCGCTG ATCCGATTTC
 H39840 CAGT.CT.GG CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ATC.GATTTC
 H95233 CAGTTCT..G CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ATC.GATTTC
 H39841 CAGT.CT.GG CAGCAG.CCC CGAGTTGTTN .CC.TCGCTG ATC.GATNTC
 N30199 CAGT.CT.GG CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ATC.GATTTC
 T52966 CAGT.CT.GG CAGCAG..CC CGAGTTGTTT .CC.TCGCTG ATG.GATTTC
 N29508 CAGT.CT..G CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ATC.GATTTC
 NZ6919 CAGT.CTTGG CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ANC.CATTTC
 N26910 CAGT.CT..G CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ATCGGATTTC
 H16757 CAGINGT.GG CAGCAGACCC CGAGITGITT .CC.TCGCTG ATC.GATTTC
N27732 CAGT.CT.GG CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ANC.GATTTC
        1451
                                                         1500
Bikunin TTT CCTCCA GGTAG AGT TTTC TTTG CTTATGTTGA ATTCCATTGC
R73968 TTTTCCTCCA GGTAAGAATT TTTCTTTT
H39840 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC
H95233 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC
H39841 TIT.CCCCCA GGTAG..AGT TITC.TITG. CITATGTTGA ANTCCATTGC
```

R73968 TTTTCCTCCA GGTAG AGT TTTC TTTG CTTATGTTGA ATTCCATTGC R73968 TTTTCCTCCA GGTAGAATT TTTCTTT
H39840 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC H95233 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC H39841 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC H39841 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC T52966 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC N29508 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC N26919 TTT.CCNCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC N26910 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC N26910 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC N27932 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC N27732 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC N27732 TTT.CCTCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ATTCCATTGC

```
Figure 4C (Con't)
          1501
                                                           1550
 Bikunin CICITI CT CATCACAGAA STSATSITSS AATCSITTET TITSTIT ST
 H39840 CICITIT.CI CATCACAGAA SISAISIISS AATGSIITSI TITSIITIISI
 H95233 CTCTTTT.CT CATCACAGAA GTGATGTTGG AATCGTTTGT TTTGTTT.GT
 H39841 STCTTTLST CATCACAGAA STSATSTISG AATCGTTTST TITGTTT.ST
 N30199 CTCTTTT.CT CATCACAGAA GTGATGTTGG AATCGTTTGT TTTGTTT.GT
 TS2966 CTCTTTT.CT CATCACAGAA GTGATGTTGG AATCGTTTGT TTTGTTT.GT
 N29508 CICITIT OF CATCACAGAA GTGATGTTGG AATCGTTTCT TITGTTT.GT
 N26919 CTCTTTT.ON CATCACAGAA GTSATGTTSG AATCGTTTCT TTTGTTT.GT
 N26910 CTCTTTT.CT CATCACAGAA GTGATGTTGG AATCGTTTCT TTTGTTT.GT
 #16757 CTCTTTTACT CATCACAGAA GTGATGTTGG AATCGTTTCT TTIGTTT.GT
 827732 CTCTTTT.CT CATCACAGAA GTGATGTTGG AATGGTTTGT TTTGTTT.GT
         1551
                                                           1600
Bikunin CTGATTTATG G TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 H39840 CTGATTTATG GGTTTTTTT AAGTAT
 H95233 CTGATTTATG G..TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 H39841 CTGATTTATG G...TTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 N30199 CTGATTTATG G. . TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 152966 CTGATTTATG G. . TITTTTT AAGTATAAC AAAAGTTTTT TATTAGCATT
 N29508 CTGATTTATG G..TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 926919 CTGATTTATG G.. ITTTTTT AAGTNTAAAC AAAAGTTTTT TATTAGCATT
 N26910 CTGATTTATG G..TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 H16757 CTGATITATG G.. TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 N27732 CTGATTTATG G. . TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
        1601
Bikunin CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
 H95233 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAA
H39841 CTGAAAGAAG GAAAGTAAAN TGTACAAGTT TAATAAAAAG GGGCCTTCCC
N30199 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
 T52966 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
N29508 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
N26919 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
N26910 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
H16757 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
N27732 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
        1651
```

N27732 CTTTAG. AAT AAAAAAAAA AAAAAAAAA AAAAAAAAA

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### FIGURE 4D

EST	consens	MLRAEADGVS	RLLGSLLLSG	VLAADRERSI	HDFCLVSKVV	GRCRASMPRW	50
EST	consens	WYNVTDGSCQ	LFVYGGCDGN	SNNYLTKEEC	LKKCATVTEN	ATGDLATSRN	100
EST	consens	AADSSVPSAP	RRQDSEDHSS	DMFNYEEYCT	ANAVTGPCRA	SFPRWYFDVE	150
EST	consens	RNSCNNFIYG	GCRGNKNSYR	SEEACMLRCF	RQQENPPLPL	GSK <u>YYYLAGL</u>	200
EST	consens	EVMVLILELG	<u>ASMVYLI</u> RVA	RRNGERALRT	VWSSGDDKEQ	LVKNTYVL	248

### FIGURE 4E

cDNA translation	ACC T	
cDNA translation	TGATCGCGAG ACCCCAACGG CTGGTGGCGT CGCCTGCGCG TCTCGGCTGA	53 -30
cDNA translation	GCTGGCCATG GCGCAGCTGT GCGGGCTGAG GCGGAGCCGG GCGTTTCTCG	103 -13
cD <b>NA</b> translation	CCCTGCTGGG ATCGCTGCTC CTCTCTGGGG TCCTGGCGGC CGACCGAGAA :	153
cDNA translation	CGCAGCATCC ACGACTTCTG CCTGGTGTCG AAGGTGGTGG GCAGATGCCG 2 1 R S I H D F C L V S K V V G R C R	203
cDNA translation	GGCCTCCATG CCTAGGTGGT GGTACAATGT CACTGACGGA TCCTGCCAGC 2 A S M P R W W Y N V T D G S C Q L	?53 38
cDNA translation	TGTTTGTGTA TGGGGGCTGT GACGGAAACA GCAATAATTA CCTGACCAAG 3 F V Y G G C D G N S N N Y L T K	303 54
cD <b>NA</b> tr <b>anslat</b> ion	GAGGAGTGCC TCAAGAAATG TGCCACTGTC ACAGAGAATG CCACGGGTGA 3 E E C L K K C A T V T E N A T G D	153 71
cD <b>NA</b> translation	CCTGGCCACC AGCAGGAATG CAGCGGATTC CTCTGTCCCA AGTGCTCCCA 4 L A T S R N A A D S S V P S A P R	03
cD <b>NA</b> tr <b>ansla</b> tion	GAAGGCAGGA TTCTGAAGAC CACTCCAGCG ATATGTTCAA CTATGAAGAA 4 R Q D S E D H S S D M F N Y E E 1	53 04
oDNA translation	TACTGCACCG CCAACGCAGT CACTGGGCCT TGCCGTGCAT CCTTCCCACG 5 Y C T A N A V T G P C R A S F P R 1	03 21
DNA translation	CTGGTACTTT GACGTGGAGA GGAACTCCTG CAATAACTTC ATCTATGGAG 5 W Y F D V E R N S C N N F I Y G G 1	53 38
DNA ranslation	GCTGCCGGGG CAATAAGAAC AGCTACCGCT CTGAGGAGGC CTGCATGCTC 6 C R G N K N S Y R S E E A C M L 1	03 54
DNA ranslation	CGCTGCTTCC GCCAGCAGGA GAATCCTCCC CTGCCCCTTG GCTCAAAGGT 6 R C F R Q Q E N P P L P L G S K Y 1	53 71
DNA ranslation	GGTGGTTCTG GCGGGGCTGT TCGTGATGGT GTTGATCCTC TTCCTGGGAG 7 V V L A G L F V M V L I L F L G A 1	C 3 8 8
DNA ranslation	CCTCCATGGT CTACCTGATC CGGGTGGCAC GGAGGAACCA GGAGCGTGCC 7 S M V Y L I R V A R R N Q E R A 2	53 04
		82 13

#### FIGURE 4F

cDNA cDNA cDNA cDNA cDNA cDNA cDNA cDNA	GCACGAGTTG GGAGGTGTAG CGCGGCTCTG AACGCGCTGA GGGCCGTTGA 50 GTGTCGCAGG CGGCGAGGGC GCGAGTGAGG AGCAGACCCA GGCATCGCGC 100 GCCGAGAAGG CCGGGGGTCC CCACACTGAA GGTCCGGAAA GGCGACTTCC 150 GGGGGCTTTG GCACCTGGCG GACCCTCCCG GAGCGTCGGC ACCTGAACGC 200 GAGGCGCTCC ATTGCGCGTG CGCGTTGAGG GGCTTCCCGC ACCTGATCGC 250 GAGACCCCAA CGGCTGGTGG CGTCGCCTGC GCGTCTCGGC TGAGCTGGCC 300 ATGGCGCAGC TGTGCGGGCT GAGGCGGAGC CGGGCGTTTC TCGCCCTGCT 35 M A Q L C G L R R S R A F L A L L -11	
cDNA translation	GGGATCGCTG CTCCTCTCTG GGGTCCTGGC GGCCGACCGA GAACGCAGCA 400	•
translation	TCCACGACTT CTGCCTGGTG TCGAAGGTGG TGGGCAGATG CCGGGCCTCC 450	
cDNA translation	ATGCCTAGGT GGTGGTACAA TGTCACTGAC GGATCCTGCC AGCTGTTTGT 500	
cDNA translation	GTATGGGGGC TGTGACGGAA ACAGCAATAA TTACCTGACC AAGGAGGAGT 550 Y G G C D G N S N N Y L T K E E C 57	
cDNA translation	GCCTCAAGAA ATGTGCCACT GTCACAGAGA ATGCCACGGG TGACCTGGCC 600 L K K C A T V T E N A T G D L A 73	
cDNA translation	ACCAGCAGGA ATGCAGCGGA TTCCTCTGTC CCAAGTGCTC CCAGAAGGCA 650 T S R N A A D S S V P S A P R R Q 90	
cDNA translation	GGATTCTGAA GACCACTCCA GCGATATGTT CAACTATGAA GAATACTGCA 700 D S E D H S S D M F N Y E E Y C T 107	
cDNA translation	CCGCCAACGC AGTCACTGGG CCTTGCCGTG CATCCTTCCC ACGCTGGTAC 750 A N A V T G P C R A S F P R W Y 123	
cDNA translation	TTTGACGTGG AGAGGAACTC CTGCAATAAC TTCATCTATG GAGGCTGCCG 800 F D V E R N S C N N F I Y G G C R 140	
cDNA translation	GGGCAATAAG AACAGCTACC GCTCTGAGGA GGCCTGCATG CTCCGCTGCT 850 G N K N S Y R S E E A C M L R C F 157	
cDNA translation	TCCGCCAGCA GGAGAATCCT CCCCTGCCCC TTGGCTCAAA GGTGGTGGTT 900 R Q Q E N P P L P L G S K <u>Y Y Y</u> 173	
cDNA translation	CTGGCGGGC TGTTCGTGAT GGTGTTGATC CTCTTCCTGG GAGCCTCCAT 950 L A G L F V M V L I L F L G A S M 190	
cDNA translation	GGTCTACCTG ATCCGGGTGG CACGGAGGAA CCAGGAGCGT GCCCTGCGCA 1000 V Y L I R V A R R N Q E R A L R T 207	}
~~NA	COGTOTGGAG OTOGGAGAT GAGAAGGAGO AGOTGGTGAA GAAGAGATAT 1050	

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### FIGURE 4F (Con't)

CDNA	ATGTGTGAGC	TTTTTTTAAA	TAGAGGGATT	GACTCGGATT	TGAGTGATCA	1150
CDNA	TTAGGGCTGA	GGTCTGTTTC	TCTGGGAGGT	AGGACGGCTG	CTTCCTGGTC	1200
CDNA	TGGCAGGGAT	GGGTTTGCTT	TGGAAATCCT	CTAGGAGGCT	CCTCCTCGCA	1250
CDNA	TGGCCTGCAG	TCTGGCAGCA	GCCCCGAGTT	GTTTCCTCGC	TGATCGATTT	1300
CDNA	Clitccicca	GGTAGAGTTT	TCTTTGCTTA	TGTTGAATTC	CATTGCCTCC	1350
CDNA	TTTTCTCNAT	CACAGAAGTG	ATGTTGGAAT	CGTTTCTTTT	GTTTGTCTGA	1400
CDNA	TTTATGGTTT	TTTTAAGTAT	AAACAAAAGT	TTTTTATTAG	CATTCTGAAA	1450
CDNA	GAAGGAAAGT	<b>AAAATGTACA</b>	AGTTTAATAA	AAAGGGGCCT	TCCCCTTTAG	1500
CDNA	AATAAATTTC	CAGCATGTTG	CTTTCAAAAA		AAAA	
1550						

#### FIGURE 4G

ESI consens			MLR	AEADGVSRLL	GSLLLSGVLA	-
PCR clone			MAQLCGL	RRSRAFLALL	GSLLLSGVLA	
AcDNA clone			Maglegl	RRSRAFLALL	GSLLLSGVIA	- 3
EST consens	ADRERSIHDE	CLVSKVVGRC	RASMPRWWYN	VTDGSCOLEV	YCCCDCNEND:	. م
r Cr CIOne	WE KE KOT UDE	CLVSKVVGRC	RASMPREMYN	UTDOSCOLER	VCCC00 011010	
AcDNA clone	ADRERSINDE	CLVSKVVGRC	RASMPRWEVE	VIDGSCQLEV	IGGCDGNSNN	5 (
			INDIA KHHIM	AIDGSCOTEA	YGGCDGNSNN	50
EST consens	YLTKEECLKK	CATVTENATG	DLATSRNAAD	SSVPSAPRRO	DSEDHSSDME	100
ECK CIONE	TELLEPOLIKK	CATVTENATG	DLATSRNAAD	SSUDSADDDO	DEEDHEEDIG	
AcDNA clone	YLTKEECLKK	CATVTENATG	DLATSRNAAD	SSVPSADAD	DSEDUSSOME	100
EST consens	NYEEYCTANA	VTGPCRASEP	RWYFDVERNS	CNNETYGGCD	CNANCABCEE	
rck clone	MILLICIANA	VTGPCRASFP	RWYFDVERNS	CNNFTYGGCB	CMMNEYBEEF	1 5 0
AcDNA clone	NYEEYCTANA	VTGPCRASEP	RWYFDVERNS	CNNETYCECE	CHENCADOLL	120
EST consens .	ACMLRCFRQQ	ENPPLPLGSK	YVYLAGLEVM	VLILFLGASM	WYTTRUADDW	200
rck clone	ACMURCEROO	ENPPLPLGSK	VVVLAGLEVM	VITTETCACH	INT TRUE PRO	
λcDNA clone .	ACMLRCFROO	ENPPLATESK	WWIAGEST	VI II EL CASH	YILLKVARRN	200
		D 1 24 2001	TTTLAG_FVA	YLLLE LGASM	<u>YYLI</u> RV <b>A</b> RRN	200
EST consens (	DERALRTVWS	SGDDKFOLVK	NTYUI			
PCR clone (	DERALRIVES	FGD				225
CDNA clone (	DERALETYMS	SCHUREULAN	UTVIII			213
		PODDUSČEAK	MIIAT			225

Purification of Placental Bikunin using Superdex 75 Gel-Filtration

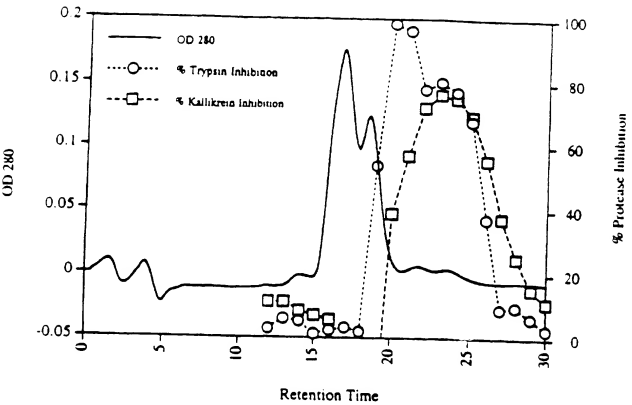
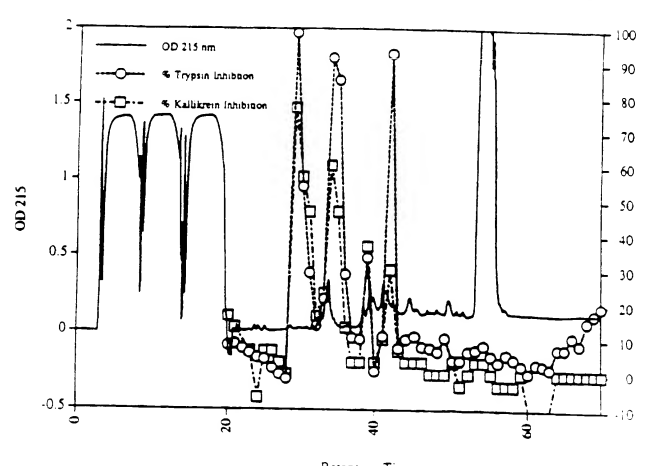


FIGURE 5

FIGURE 6

# Purification of Placental Bikunin using C18 Reverse-Phase Chromatography



Retention Time

% Protease Inhibition

Figure 7

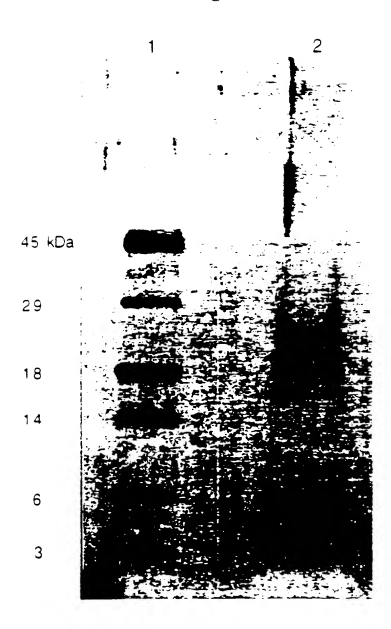


Figure 8A

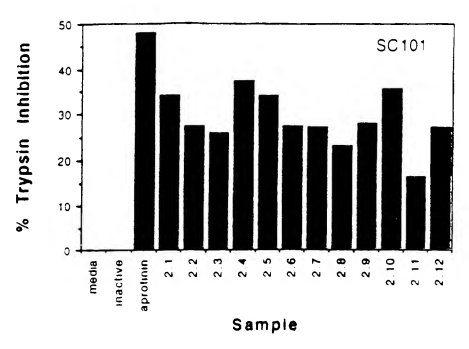


Figure 8B

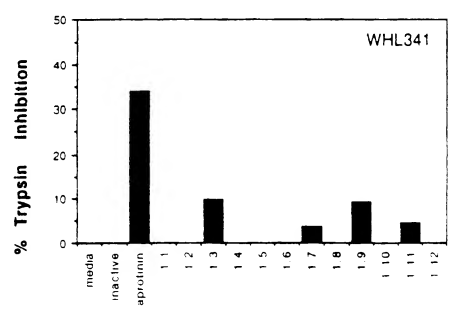


Figure 9

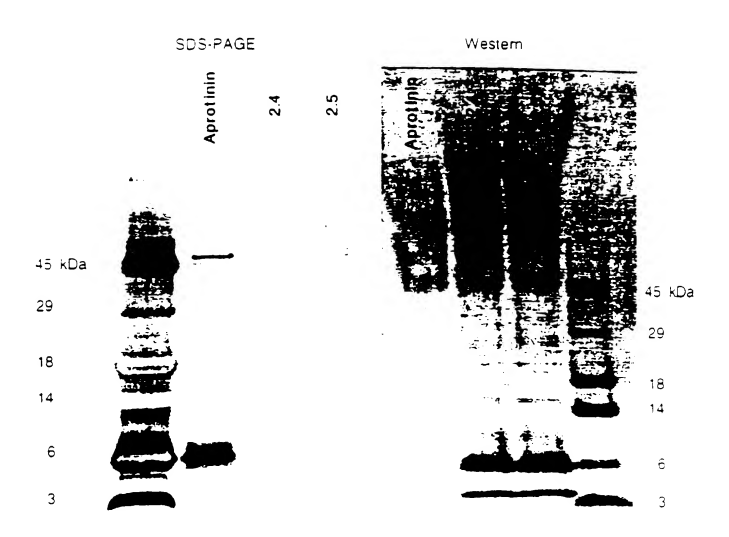
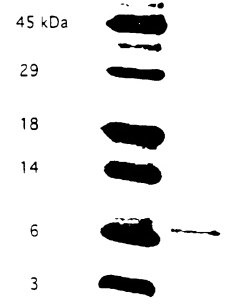


Figure 10

1 2



igure 11

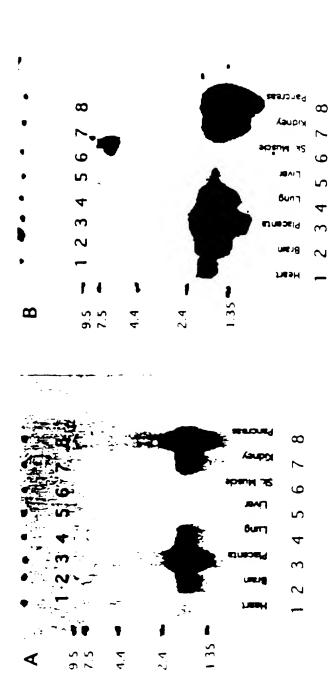


Figure 12

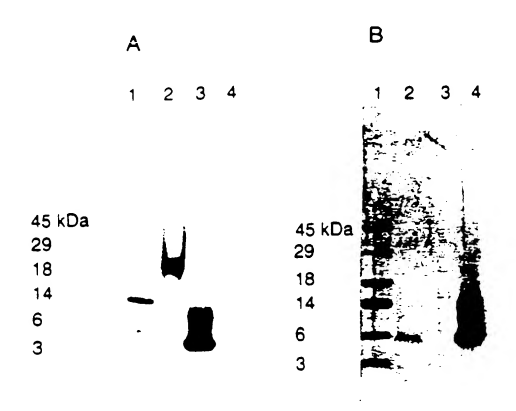


Figure 13

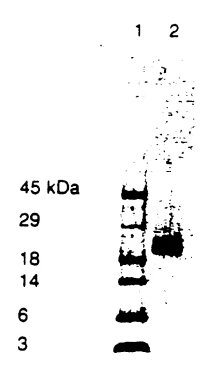
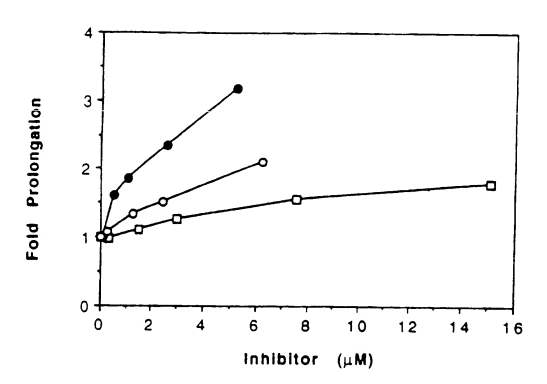
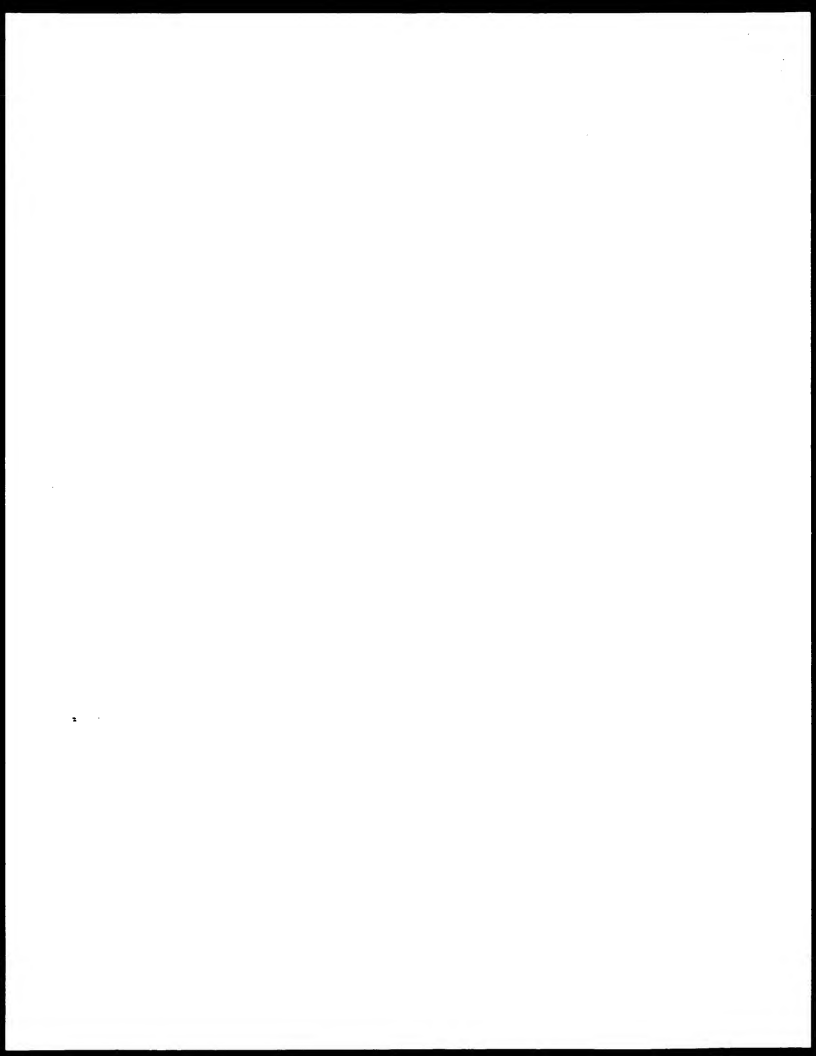


Figure 14





# **PCT**

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



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(22) International Filing (30) Priority Data:	Date: 10 March 19	97 (10.03.9°	MARLOR, Christopher, W. [US/US]. 11 Robertson Drive Bethany, CT 06524 (US). MULLER, Daniel, K. [US-US] 253 Hemlock Hill Road, Orange, CT 06477 (US).	
60/013,106 60/019,793 08/725,251	11 March 1996 (11.03.9 14 June 1996 (14.06.96) 4 October 1996 (04.10.9	) U	e. Danning 2000 South Wacker Drive ( hicago II Milhill	
(60) Parent Application: (63) Related by Cont.			(81) Designated States: AL, AM, AT, AU, AZ, BB, BG, BR, BY CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD	
US Filed on US Filed on US Filed on US Filed on	4 October 19 60/0 14 June 19	)19,793 (CII 96 (14.06.90 )13,106 (CII	SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN ARIPO patent (GH, KE, LS, MW, SD, SZ, UG), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT)	

CORPORATION [US/US]; One Mellon Center, 500 Grant Street, Pittsburgh, PA 15219-2507 (US).

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(75) Inventors/Applicants (for US only): TAMBURINI, Paul, P. [GB/US]; 36 Misty Mountain Road, Kensington, CT 06460 With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(88) Date of publication of the international search report: 13 November 1997 (13 11 97)

## (54) Title: HUMAN BIKUNIN

#### (57) Abstract

The instant invention provides for proteins, polypepudes, nucleic acid sequences, constructs, expression vectors, host cells, pharmaceutical compositions of, and methods for using human placental bikunin, serine protease inhibitor domains, and fragments thereof.

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lnten nal Application No PCT/US 97/03894

A. CLASSIFICATION OF SUBJECT MATTER C07K14/81 A61K38/57 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) 1PC 6 C07K Documentation searched other than manimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X EMBL/GENBANK DATABASES Accession no R35464 1-6,11 Sequence reference HS46499, May 4, 1995 L.HILLIER ET AL: "The WashU-Merck EST Project\* XP002039653 see the whole document X EMBL/GENBANK DATABASES Accession no N39798 1-6.11Sequence reference HS798277, January 26, L. HILLIER ET AL: "The Wasu-Merck EST Project\* XP002039654 see the whole document -/--X Further documents are listed in the continuation of box C. X Patent family members are listed in annex. Special categories of cited documents: T later document published after the international filing date or priority date and not in conflict with the applicati 'A' document defining the general state of the art which is not considered to be of particular relevance. ated to understand the principle or theory underlying the invention cartier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date and set the authors and first SLACK . II AG R. & When special reason has specified AUTHOR THE GETTING GETTER OF THE WIND AND THE STATE OF TH യ്ക്കുന്നു പ്രധാന വേശ്യാവരുന്നു. വ Kher means nents, such combination being obvious to a person still cu document published prior to the international filing date but later than the priority date claimed "A" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report n 1, 18, 97

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European Patent Office, P.B. 5818 Patendaan 2. NL: 2280 HV Ripmith Ta 1 + 3 TO) 340-2040. Tx 3, 411 cpn nt

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Category '	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
X	EMBL/GENBANK DATABASES Accession no R74593 Sequence reference HS593137, June 9, 1995 L.HILLIER ET AL: "The WashU-Merck EST Project" XP002039655 see the whole document	1-6,11
P,X	EP 0 758 682 A (MITSUBISHI CHEM CORP) 19 February 1997 see the whole document	1-11
P,A	JOURNAL OF BIOLOGICAL CHEMISTRY 272 (10). 1997. 6370-6376. ISSN: 0021-9258, XP002039700 SHIMOMURA T ET AL: "Hepatocyte growth factor activator inhibitor, a novel Kunitz-type serine protease inhibitor."	
A	DATABASE MEDLINE accession no 94289695, 1 July 1994 J. WOJTA ET AL: "Hepatocyte growth factor stimulates expression of plasminogen activator inhibitor type 1 and tissue factor in HepG2 cells" XP002039702 see abstract & BL00D, vol. 84, no. 1, 1994, pages 151-157,	7-10
A	JOURNAL OF BIOLOGICAL CHEMISTRY 271 (7). 1996. 3615-3618. ISSN: 0021-9258, XP002039701 MIYAZAWA K ET AL: "Activation of hepatocyte growth factor in the injured tissues is mediated by hepatocyte growth factor activator."	7-10

Intrimetional application No.

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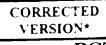
Box I Observations where certain claims were found unsearchable (Continuation of Iter	n 1 of first sheet)
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Claims Nos because they relate to subject matter not required to be searched by this Authority, namely Remark: Although claim(s) 7-9 is(are) directed to a method of treatment of the h body, the search has been carried out and based on effects of the compound/composition.  Claims Nos because they relate to parts of the International Application that do not comply with the prescribed in an extent that no meaningful international Search can be carried out, specifically	the alleged
Claims Nos because they are dependent claims and are not drafted in accordance with the second and third se	
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This International Searching Authority found multiple inventions in this international application, as follows	
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As all searchable claims could be searched without effort justifying an additional fee, this Authority of any additional fee.	did not invite payment
As only some of the required additional search fees were timely paid by the applicant, this internation covers only those claims for which fees were paid, specifically claims Nos	ona: Search Report
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Patent document cated in search report	Publication date	Patent family member(s)	Publication date
EP 0758682 A	19-02-97	JP 9095498 A	08-04-97



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# REATY (PCT)

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(21) International Application Number: PCT US( (22) International Filing Date: 10 March 1997 ( (30) Priority Data: 60:013,106 11 March 1996 (11.03.96) 60:019.793 14 June 1996 (14.06.96) 08:725,251 4 October 1996 (04.10.96)	10.03.9 L	ford, CT 06460 (US). DELARIA, Katherine, A. [US US].			
(60) Parent Applications or Grants (63) Related by Continuation US Filed on	04.10.9 793 (CD 14.06.9 06 (CD	SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, ARIPO patent (GH, KE, LS, MW, SD, SZ, UG), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT,			

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(54) Title: HUMAN BIKUNIN

(57) Abstract

The instant invention provides for proteins, polypeptides, nucleic acid sequences, constructs, expression vectors, host cells, pharmaceutical compositions of, and methods for using human placental bikunin, serine protease inhibitor domains, and fragments thereof

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#### Human Bikunin

#### Field of the Invention

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The compositions of the invention relate to the field of proteins which inhibit serine protease activity. The invention also relates to the field of nucleic acid constructs, vectors and host cells for producing serine protease inhibiting proteins, pharmaceutical compositions containing the protein, and methods for their use.

## 10 Background of the Invention

#### Problem Addressed

Blood loss is a serious complication of major surgeries such as open heart surgery and other complicated procedures. Cardiac surgery patients account for a significant proportion of transfused donor blood. Blood transfusion carries risks of disease transmission and adverse reactions. In addition, donor blood is expensive and demands often exceed supply. Pharmacological methods for reducing blood loss and the resultant need for transfusion have been described (reviewed by Scott et al., Ann. Thorac. Surg. 50: 843-851, 1990).

#### 20 Protein Serine Protease Inhibitor

Aprotinin, a bovine serine protease inhibitor of the Kunitz family is the active substance in the medicament Trasylol®. Aprotinin (Trasylol®) has been reported as being effective in reducing perioperative blood loss (Royston et al., Lancet ii: 1289-1291, 1987; Dietrich et al., Thorac. Cardiovasc. Surg. 37: 92-98 1989; Fraedrich et al., Thorac. Cardiovasc. Surg. 37: 89-91, 1989); W. van Oeveren et al. (1987), Ann Thorac. Surg. 44, pp 640-645; Bistrup et al., (1988) Lancet I, 366-367), but adverse effects, including hypotension and flushing (Bohrer et al., Anesthesia 45: 853-854, 1990) and allergic reactions (Dietrich et al., Supra) have been reported. Use of aprotinin in patients previously exposed to it is not recommended (Dietrich et al., Supra). Trasylol® has also been used for the treatment of hyperfibrinolytic hemorrhages and traumatic hemorrhagis

The motry psing prasming and rank rend and is used the rapeutically in the streament of acute pancreatitis, various states of shock syndrome by perfibring the bemorrhage and myotardial infarction (Trappell et al., 1971) and hear the size of the stream of

al., (1979)Acta Neurochir. 49: 207; Sher (1977) Am J. Obstet. Gynecol. 129: 164; Schneider (1976), Artzneim.-Firsch. 26: 1606). It is generally thought that Trasylol® reduces blood loss *in vivo* through inhibition of kallikrein and plasmin. It has been found that aprotinin (3-58, Arg15, Ala17, Ser42) exhibits improved plasma kallikrein inhibitory potency as compared to native aprotinin itself (WO 89/10374).

# Problems With Aprotinin

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Because aprotinin is of bovine origin, there is a finite risk of inducing anaphylaxis in human patients upon re-exposure to the drug. Thus, a human functional equivalent to aprotinin, by virtue of a lower risk of anaphylaxis, would be most useful and desirable to have.

Aprotinin is also nephrotoxic in rodents and dogs when administered repeatedly at high dose (Bayer, Trasylol<sup>®</sup>, Inhibitor of proteinase; Glasser et al., in "Verhandlungen der Deutschen Gesellschaft für Innere Medizin, 78. Kongress", Bergmann, Munchen, 1972 pp. 1612-1614). One hypothesis ascribes this effect to the accumulation of aprotinin in the negatively charged proximal tubules of the kidney, due to its high net positive charge (WO 93/14120).

Accordingly, an object of the present invention is to identify human proteins with functional activity similar to aprotinin. It was also an object of the instant invention to identify human proteins, that would be less charged, yet exhibit the same, highly similar, or improved protease specificities as found for aprotinin, especially with respect to the potency of plasmin and kallikrein inhibition. Such inhibitors could then be used repeatedly as medicaments in human patients with reduced risk of adverse immune response and reduced nephrotoxicity.

# Brief Summary of the Invention

The instant invention provides for a purified human serine protease inhibitor which can specifically inhibit kallikrein, that has been isolated from human placental tissue via affinity chromatography.

The instant invention provides a newly identified human protein herein called human placental bikunin that contains two serine protease inhibitor domains of the Kunitz class. In one particular embodiment, the instant invention embodies a protein having the amino acid sequence:

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ADFERSIHDE DIVSKVVGED RASMEEWWYN VTDGSDQLEV YGGOLGNSNN: 8.
YLTKEECLKK DATVTENATG DIATSENAAD GSVESAFREQ DSEDHSSDMF 10.0
NYEEYOTANA VTGECRASEE RWYFDVERNS DIMFIYGGOF GNKNSYRSEE 15.0
ADMIRGERQQ ENEELPLGSK VTVLAGAVS
```

In a prefered embodiment the instant invention provides for native human placental bikunin protein having the amino acid sequence:

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ADRERSIHDE GLVSKVVGRG RASMPRWWYN VTDGSGQLEV YGGGDGNSNN: 50
YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDME 100
NYEEYGTANA VTGECRASEP RWYEDVERNS ONNEIYGGGR GNKNSYRSEE 150
ACMLRGERQQ ENEPLPLGSK 171
(SEQID NO:52)
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WO 97 33996

(SEQ ID NO: 1)

In one aspect, the biological activity of the protein of the instant invention is that it can bind to and substantially inhibit the biological activity of trypsin, human plasma and tissue kallikreins, human plasmin and Factor XIIa. In a preferred embodiment, the present invention provides for a native human placental bikunin protein, in glycosylated form. In a further embodiment the instant invention encompasses native human bikunin protein which has been formed such that it contains at least one cysteine-cysteine disulfide bond. In a preferred embodiment, the protein contains at least one intra-chain cysteinecysteine disulfide bond formed between a pair of cysteines selected from the group consisting of CYS11-CYS61, CYS20-CYS44, CYS36-CYS57, CYS10n-CYS156, CYS115-CYS139, and CYS131-CYS152, wherein the cysteines are numbered according to the amino acid sequence of native human placental bikunin. One of ordinary skill will recognize that the protein of the instant invention may fold into the proper three-dimensional conformation, such that the biological activity of native human bikurin is maintained, where none, one or more, or all of the native intra-chain cysteine-cysteine disulfide bonds are present. In a most preferred embodiment, the protein of the instant invention is

Active protein of the instant invention can be obtained by purification from human tissue, such as placenta, or via synthetic protein chemistry to the passes a strate file the Evangles release. It is used to restore the true

protein of the instant invention may be obtained using molecular biology techniques, where self-replicating vectors are capable of expressing the protein of the instant invention from transformed cells. Such protein can be made as non-secreted, or secreted forms from transformed cells. In order to facilitate secretion from transformed cells, to enhance the functional stability of the translated protein, or to aid folding of the bikunin protein, certain signal peptide sequences may be added to the NH2-terminal portion of the native human bikunin protein.

In one embodiment, the instant invention thus provides for the native human bikunin protein with at least a portion of the native signal peptide sequence intact. Thus one embodiment of the invention provides for native human bikunin with at least part of the signal peptide, having the amino acid sequence:

15 AGSFLAWLGSLLLSGVLA -1
ADRERSIHDFCLVSKVVGRCRASMPRWWYNVTDGSCQLFVYGGCDGNSNN 50
YLTKEECLKKCATVTENATGDLATSRNAADSSVPSAPRRQDSEDHSSDMF 100
NYEEYCTANAVTGPCRASFPRWYFDVERNSCNNFIYGGCRGNKNSYRSEE 150
ACMLRCFRQQENPPLPLGSKVVVLAGAVS 179
20 (SEQ ID NO: 2)

In a prefered embodiment the instant invention provides for a native human placental bikunin protein with part of the leader sequence intact, having the amino acid sequence of SEQ ID NO: 52 with an intact leader segment having the amino acid sequence:

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MAQLCGL RRSRAFLALL GSLLLSGVLA -1 (SEQ ID NO: 53)
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In another embodiment, the instant invention provides for bikunin protein with part of the leader sequence intact, having the amino acid sequence of SEQ ID NO: 52 with the intact leader segment having the amino acid sequence:

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MLR AEADGVSRLL GSLLLSGVLA -1 (SEQ ID NO: 54)
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In a preferred numbering system used herein the amino acid numbered +1 is assigned to the NH2-terminus of the amino acid sequence for native

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human placental bikunin. One will readily recognize that functional protein fragments can be derived from native human placental bikunin, which will maintain at least part of the biological activity of native human placental bikunin, and act as serine protease inhibitors.

In one embodiment, the protein of the instant invention comprises a fragment of native human placental bikunin, which contains at least one functional Kunitz-like domain, having the amino acid sequence of native human placental bikunin amino acids 7-159, hereinafter called "bikunin (7-159)". Thus the instant invention embodies a protein having the amino acid sequence:

```
IHDFOLVSKVVGRORASMFFWWYNVTDGSOQLFVYGGODGNSNN 50
YLTKEEOLKKOATVTENATGDLATSRNAADSSVFSAFRRQDSEDHSSDMF 100
NYEEYOTANAVTGFORASFPRWYFDVERNSONNFIYGGORGNKNSYFSEE 160
ACMLROFFQ 159
(SEQ ID NO:3)
```

where the amino acid numbering corresponds to that of the amino acid sequence of native human placental bikunin. Another functional variant of this embodiment can be the fragment of native human placental bikunin, which contains at least one functional Kunitz-like domain, having the amino acid sequence of native human placental bikunin amino acids 11-156, bikunin (11-156)

```
25 OLUSKUUGE ORASMEE WWYNDUTOGGOQLEUY GGOOGNISMI SUU
YLTKEEOLKKOATUTENATGOLATSENAADISSUPSAFREQOISEDHISOME 1
NYEEYOTANAUTGEOFASEPEWYFOUERNISOMEFTY GGOFGNENISYESEE 15
ACMLRO (SEQ ID NO: 50).
```

One can recognize that the individual Kunitz-like domains are also

and the second second

bikunin amino acids 7-64, hereinatter called "bikunin (7-64)". Thus in one embodiment the instant invention encompasses a protein which contains at the local factors of the protein embodiment.

IHDFOLVSHIVIGFORASMFRWWYNVTDGSOQLFUYGGODGNSINI (50 YLTKEEDLKKOATU 64 (SEQ ID NO:4)

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where the amino acid numbering corresponds to that of the amino acid sequence of native human placental bikurun. Another form of the protein of the instant invention can be a first Kunitz-like domain consisting of the amino acid sequence of native human placental bikurun amino acids 11-61, "bikurun (11-61)" having the amino acid sequence:

CLVSKVVGRCRASMPRWWYNVTDGSCQLFVYGGCDGNSNN 50 YLTKEECLKKC 51 (SEQ ID NO: 5)

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The instant invention also provides for a protein having the amino acid sequence of a Kunitz-like domain consisting of the amino acid sequence of native human placental bikunin amino acids 102-159, hereinafter called "bikunin (102-159)". Thus one embodiment the instant invention encompasses a protein which contains at least one Kunitz-like domain having the amino acid sequence:

YEEYOTANAUTGPORASFPRWYFDVERNSONNFINGGORGNKNSYRSEE 150 ACMLROFRQ 159 (SEQ ID NO: 6)

where the amino acid numbering corresponds to that of the amino acid sequence of native human placental bikunin. Another form of this domain can be a Kunitz-like domain consisting of the amino acid sequence of native human placental bikunin amino acids 106-156, "bikunin (106-156)" having the amino acid sequence:

OTANAVTGFOFASFPRWYFOVERNSONNFIYGGOFGNKNSYFSEE 15.
ACMLRC 156
(SEQIDNO:7)

Thus one of ordinary skill will recognize that fragments of the native

human bikunin protein can be made which will retain at least some of the native protein biological activity. Such fragments can also be combined in different orientations or multiple combinations to provide for alternative proteins which retain some of, the same, or more biological activity of the native human bikunin protein.

One will readily recognize that biologically active protein of the instant invention may comprise one or more of the instant Kunitz-like domains in combination with additional Kunitz-like domains from other sources. Biologically active protein of the instant invention may comprise one or more of the instant Kunitz-like domains in combination with additional protein domains from other sources with a variety of biological activities. The biological activity of the protein of the instant invention can be combined with that of other known protein or proteins to provide for multifunctional fusion proteins having predictable biological activity. Thus, in one embodiment, the instant invention encompasses a protein which contains at least one amino acid sequence segment the same as, or functionally equivalent to the amino acid sequence of either SEQ ID NO: 5 or SEQ ID NO: 7.

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An open reading frame which terminates at an early stop codon can still code for a functional protein. The instant invention encompasses such alternative termination, and in one embodiment provides for a protein of the amino acid sequence:

ADRERSIHDFOLVSKVVGROFASMFRWWYNVTDGSOQLFVVGGGOGNSND: 50 YLTKEEOLKKOATVTENATGOLATSFNAADSSVFSAPFFQDS 30 (SEQIDNO:8)

In one embodiment, the instant invention provides for substantially purified, or recombinantly produced native human bikunin protein with an intact segment of the leader sequence, and at least a portion of the native transmembrane region intact. Thus one embodiment of the invention provides for native human bikunin, with an intact leader sequence, and with at least part

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MER ABADOVERUL SELLESOVIA
     1 E3T
                              MAQLOGL FRSFAFLALL GSLULGGVLA,
     J POF
                              MAQLOGL RESEAFLALL GSLLLSGVLA
    3)À cDNA
5
     1 ADRERSINGE CLUSHUNGES PASMPRWAYN UTDGSSQLEN YGGCDGNSMN 51
     E) ADREFSIHOF CLYSKY/GRO RASMPRWYYN UTDGSCOLFY YGGCDGNSMI 50
     3) ADREPSINGE CLUSKUUGEC FASMPRWAYN UTDGSCOLFU YGGCDGNSNN 50
     1-YLTREECLFY CATUTENATS CLATSPNAAD SEVPSAFRRQ DSECHSSOMF 100
     DYMITKEECLYK CATUTENATG DLATSENAAD SSUPSAPRRQ DSEDHSSDMF 100
10
     3)YLTKEECLFK CATTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF 100
     1) NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE 150
     2) NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE 150
     3) NYEEYOTANA UTGPORASFP RWYFDVERNS CHNFIYGGOR GHKNSYRSEE 151
15
     1) ACMERCFRQQ ENPPEPEGSK WWWLAGLFWM WEILFEGASM WYLIRWARRN 200
     2) ACMLROFRQQ ENPPLPLGSK VVVLAGLFVM VLILFLGASM VYLIRVARRN 200
     3) ACMLROFRQQ ENPPLPLGSK VVVLAGLFVM VLILFLGASM VYLIRVARRN 200
20
                                                                225
     1) QERALPTVWS SGDDKEQLVK NTYVL
     2:QERALRITANS FGD
     3 OPERALFTIAS SOCCHEQUIA NTEVIL
                                                               22E
```

where sequence 1) is EST derived consensus SEQ ID NO: 45, 2) is PCR clone 25 SEQ ID NO:47, and 3) is lambda cDNA clone SEQ ID NO:49. In a preferred embodiment a protein of the instant invention comprises one of the amino acid sequence of SEQ ID NO: 45, 47 or 49 wherein the protein has been cleaved in the region between the end of the last Kunitz domain and the transmembrane 30 region.

The instant invention also embodies the protein wherein the signal peptide is deleted. Thus the instant invention provides for a protein having the amino acid sequence of SEQ ID NO: 52 continuous with a transmembrane amino acid sequence:

35	EST	WWWLAGLFUM VLILFLGASM WYLIRWARRN	200
	EST	QERALRTVWS SGDDKEQLUK NTYVL	225
		(SEQ ID NO: 69)	
	a transmemb	orane amino acid sequence:	
	PCR	WWWLAGLFUM VLILFLGASM WYLIRVARRN	200
40	PCR	QERALRITUMS FGD	213
		(SEQ ID NO: 68)	
	or a transme	mbrane amino acid sequence:	
	<b>ACDNA</b>	WWWLAGLFUM VLILFLGASM WYLIRVAREN	200
	λcDNA	ÇERALRIYWS SGDDKEÇLUK NIYVL	225
45		(SEQ ID NO: 67).	

The protein amino acid sequences of the instant invention clearly teach one of the art the appropriate nucleic acid sequences which can be used in molecular biology techniques to produce the proteins of the instant invention. Thus, one embodiment of the instant invention provides for a nucleic acid sequence which encodes for a human bikurin having the consensus DNA sequence of Figure 3 (SEQ ID NO: 9), which translates into the amino acid sequence for native human placental bikurin sequence of Figure 3 (SEQ ID NO: 10). In another embodiment, the instant invention provides for a consensus nucleic acid sequence of Figure 4C (SEQ ID NO: 51) which encodes for an amino acid sequence of Figure 4D (SEQ ID NO: 45).

In a preferred embodiment, the instant invention provides for a nucleic acid sequence which encodes for native human placental bikunin having the DNA sequence of Figure 4F (SEQ ID NO: 48) which encodes for the protein sequence of SEQ ID NO: 49. In an another embodiment, the instant invention provides for a nucleic acid sequence of Figure 4E (SEQ ID NO: 46) which encodes for a protein sequence of SEQ ID NO: 47.

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One can easily recognize that certain allelic mutations, and conservative substitutions made in the nucleic acid sequence can be made which will still result in a protein amino acid sequence encompassed by the instant invention. One of skill in the art can recognize that certain natural allelic mutations of the protein of the instant invention, and conservative substitutions of amino acids in the protein of the instant invention will not significantly alter the biological activity of the protein, and are encompassed by the instant invention.

The instant invention also provides for pharmaceutical compositions containing human placental bikurin and fragments thereof that are useful for the reduction of perioperative blood loss in a patient undergoing surgery

The present invention also provides methods for reducing perioperative blood loss in a patient undergoing surgery, wherein an effective amount of the disclosed human serine protease inhibitors of the present invention in a biologically compatible vehicle is administered to the patient.

The present invention also provides for variants of placental bikunin,

sequence for native placental bikunin. Substitutions at Xaa+ through Xaa+7 are also preferred for variants of bikunin (7-64), while substitutions at Xaa+7 2000 and Xaa+7 2

Thus the present invention embodies protein having an amino acid sequence:

	Ala Asp Arg Ol. Arg Ser Ile Waa Asp Pne	10
	Cys Leu Val Ser Lys Val Kaa <sup>l</sup> Oly Kaa <sup>l</sup> Cys	2.3
5	Kaa <sup>4</sup> Xaa <sup>5</sup> Xaa <sup>6</sup> Xaa <sup>7</sup> Xaa <sup>8</sup> Kaa <sup>9</sup> Trp Trp Tyr Asn	3.0
	Wal Thr Asp Gly Ser Cys Gln Leu Phe Kaa <sup>l)</sup>	40
	Tyr Xaa $^{11}$ Gly Cys Xaa $^{13}$ Xaa $^{13}$ Xaa $^{14}$ Ser Asn Asn	50
	Tyr Xaa <sup>15</sup> Thr Lys Glu Glu Cys Leu Lys Lys	60
	Cys Ala Thr Xaa <sup>lo</sup> Thr Glu Asn Ala Thr Gly	~ :
10	Asp Leu Ser Thr Ser Arg Ash Ala Ala Asp	3.0
	Ser Ser Val Pro Ser Ala Pro Arg Arg Gin	9.0
	Asp Ser Glu His Asp Ser Ser Asp Met Phe	100
	Asn Tyr Maa <sup>17</sup> Glu Tyr Cys Thr Ala Asn Ala	110
	Mal Maa $^{18}$ Oly Maa $^{19}$ Cys Maa $^{20}$ Maa $^{21}$ Maa $^{22}$ Maa $^{23}$ Maa $^{24}$	12.
15	Kaa <sup>05</sup> Trp Tyr Phe Asp Val Glu Arg Ash Ser	130
	Cys Asn Asn Phe Maa <sup>26</sup> Tyr Maa <sup>27</sup> Gly Cys Maa <sup>28</sup>	140
	Xaa <sup>29</sup> Xaa <sup>33</sup> Dys Asn Ser Tyr Xaa <sup>31</sup> Ser Glu Glu	150
	Ala Cys Met Leu Arg Cys Phe Arg Kaa <sup>32</sup> Gln	160
	Glu Asn Pro Pro Leu Pro Leu Gly Ser Lys	170
20	Val Val Val Leu Ala Gly Ala Val Ser	179
	(SEQ ID NO: 11).	

where  $Xaa^1$  -  $Xaa^{32}$  each independently represents a naturally occurring amino acid residue except Cys, with the proviso that at least one of the amino acid residues  $Xaa^{1}$ - $Xaa^{32}$  is different from the corresponding amino acid residue of the native sequence.

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In the present context, the term 'naturally occurring amino acid residue' is intended to indicate any one of the 20 commonly occurring amino acids, i.e., Ala, Arg, Asn, Asp, Cys, Gln, Glu, Gly, His, Ile, Leu, Lys, Met, Phe, Pro, Ser, Thr, Trp, Tvr and Val.

By substituting one or more amino acids in one or more of the positions indicated above, it may be possible to change the inhibitor specificity profile of native placental bikunin or that of the individual Kunitz-like domains, bikunin(7-64) or bikunin (102-159) so that it preferentially inhibits other serine proteases such as, but not limited to, the enzymes of the complement cascade. TF/FVIIa, FXa, thrombin, neutrophil elastase, cathepsin G or proteinase-3.

Examples of preferred variants of placental bikunin include those

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wherein  $x_{aa}$  is an amino acid residue selected from the group consisting or His. Glu, Pro. Ala, Val or Lys. in particular wherein Xaal is His or Pro, or wherein Xaa2 is an amino acid residue selected from the group consisting of Val. Thr. Asp. Pro. Arg. Tyr. Glu. Ala. Lys. in particular wherein Xaa- is Val or Thr; or wherein Xaa<sup>3</sup> is an amino acid residue selected from the group consisting of Arg, Pro, Ile, Leu, Thr. in particular wherein Xaa<sup>3</sup> is Arg or Pro; or wherein  $Xaa^4$  is an amino acid residue selected from the group consisting of Arg, Lys and Ser, Gln, in particular wherein Xaa4 is Arg or Lys; or wherein  $Xaa^{5}$  is an amino acid residue selected from the group consisting of Ala, Gly, Asp, Thr, in particular wherein Xaa<sup>5</sup> is Ala; or wherein Xaa<sup>6</sup> is an amino acid residue selected from the group consisting of Ser, Ile, Tvr, Asn, Leu, Val, Arg, Phe, in particular wherein Xaa<sup>6</sup> is Ser or Arg; or wherein Xaa<sup>7</sup> is an amino acid residue selected from the group consisting of Met, Phe, Ile, Glu, Leu. Thr and Val, in particular wherein Xaa<sup>7</sup> is Met or Ile; or wherein Xaa<sup>8</sup> is an amino acid residue selected from the group consisting of Pro, Lvs, Thr. Gln, Asn. Leu. Ser or Ile, in particular wherein Xaa<sup>8</sup> is Pro or Ile; or wherein Xaa<sup>9</sup> is an amino acid residue selected from the group consisting of Arg, Lys or Leu, in particular wherein Xaa<sup>9</sup> is Arg: or wherein Xaa<sup>10</sup> is an amino acid residue selected from the group consisting of Val, Ile. Lvs. Ala, Pro, Phe, Trp, Gln, Leu and Thr, in particular wherein  $Xaa^{10}$  is Val; or wherein  $Xaa^{11}$  is an amino acid residue selected from the group consisting of Glv, Ser and Thr, in particular wherein Xaa<sup>11</sup> is Glv; or wherein Xaa<sup>12</sup> is an amino acid residue selected from the group consisting of Asp, Arg, Glu. Leu, Gln, Gly, in particular wherein Xaa12 is Asp or Arg; or wherein Xaa13 is an amino acid residue selected from the group consisting of GIV and Ala or wherein Xaa<sup>14</sup> is an amino acid residue. selected from the group consisting of Asn or Lys; or wherein  $\lambda_{aa}^{15}$  is an aminoacid residue selected from the group consisting of Gly, Asp. Leu, Arg. Glu, Thr. Tvr, Val, and Lvs, in particular wherein Xaa 15 is Leu or Lvs; or wherein Xaa 16 is an amino acid residue selected from the group consisting of Val. Gin. Asp. Glv. Ile, Ala, Met, and Val. in particular wherein Xaa 16 is Val or Ala; or wherein  $Xaa^{17}$  is an amino acid residue selected from the group consisting of His. Giu.

Asplaced Argustic Constant and Asplaced Argustic Argustic Consisting of Argustic Configuration and Argustic Constant Constant Argustic Constant Configuration and Argustic Configuration and Argustic Configuration and Argustic Configuration C

Ser, in particular wherein Xaa<sup>20</sup> is Arg or Lys; or wherein Xaa<sup>21</sup> is an amino acid residue selected from the group consisting of Ala, Asp, Thr or Gly, in particular wherein Xaa<sup>21</sup> is Ala; or wherein Xaa<sup>22</sup> is an amino acid residue selected from the group consisting of Ser, Ile, Tyr, Asn, Leu, Val, Arg or Phe, in particular wherein Xaa<sup>22</sup> is Ser or Arg; or wherein Xaa<sup>23</sup> is an amino acid residue selected from the group consisting of Met, Phe, Ile, Glu, Leu, Thr and Val, in particular wherein  $Xaa^{23}$  is Phe or Ile; or wherein  $Xaa^{24}$  is an amino acid residue selected from the group consisting of Pro, Lys, Thr. Asn, Leu, Gln, Ser or Ile, in particular wherein Xaa<sup>24</sup> is Pro or Ile; or wherein Xaa<sup>25</sup> is an amino acid residue selected from the group consisting of Arg, Lys or Leu, in particular wherein  $Xaa^{25}$  is Arg: or wherein  $Xaa^{26}$  is an amino acid residue selected from the group consisting of Val, Ile, Lys, Leu, Ala, Pro, Phe, Gln, Trp and Thr, in particular wherein Xaa<sup>26</sup> is Val or Ile; or wherein Xaa<sup>27</sup> is an amino acid residue selected from the group consisting of Gly, Ser and Thr, in particular wherein  $Xaa^{27}$  is Gly; or wherein  $Xaa^{28}$  is an amino acid residue selected from the group consisting of Asp, Arg, Glu, Leu, Gly or Gln, in particular wherein  $Xaa^{28}$  is Arg; or wherein  $Xaa^{29}$  is an amino acid residue selected from the group consisting of Gly and Ala; or wherein Xaa<sup>30</sup> is an amino acid residue selected from the group consisting of Asn or Lys; or wherein Xaa<sup>31</sup> is an amino acid residue selected from the group consisting of Gly, Asp, Leu, Arg, Glu, Thr, Tyr, Val, and Lys, in particular wherein Xaa31 is Arg or Lys; or wherein Xaa<sup>32</sup> is an amino acid residue selected from the group consisting of Val, Gln, Asp, Glv, Ile, Ala, Met, and Thr, in particular wherein  $Xaa^{32}$  is Gln or Ala.

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### Description of the Drawings

The invention will be better understood from a consideration of the following detailed description and claims, taken in conjunction with the drawings, in which:

Figure 1 depicts the nucleotide sequence of EST R35464 (SEQ ID NO: 12) and the translation of this DNA sequence (SEQ ID NO: 13) which yielded an open reading frame with some sequence similarity to aprotinin. The translation product contains 5 of the 6 cysteines in the correct spacing that is characteristic for Kunitz-like inhibitor domains (indicated in bold). The position normally occupied by the remaining cysteine (at codon 38) contained instead a phenylalanine (indicated by an asterisk).

Figure 2 depicts the nucleotide sequence of EST R74593 (SEQ ID NO: 14),

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and the translation of this DNA sequence (SEQ ID NO: 15) which yielded an open reading frame with homology to the Kunitz class of serine protease inhibitor domains. The translation product contained 6 cysteines in the correct spacing that is characteristic for Kunitz-like inhibitor domains (indicated in bold). However, this reading frame sequence includes stop codons at codon 3 and 23.

Figure 3 depicts a deduced nucleic acid sequence of human placental bikunin (SEQ ID NO: 9) labeled "consensus" and matched with the translated protein amino acid sequence labeled "translated" (SEQ ID NO: 10). Also as comparison are shown the nucleic acid sequence for ESTs H94519 (SEQ ID NO: 16). N39798 (SEQ ID NO: 17), R74593 (SEQ ID NO: 14) and R35464 (SFQ ID NO: 12). The underlined nucleotides in the consensus sequence correspond to the site of PCR primers described in the Examples. Underlined amino acids in the translated consensus sequence are residues whose identity have been confirmed by amino acid sequencing of purified native human placental bikunin. Nucleotide and amino acid code are standard single letter code, "N" in the nucleic acid code indicates an unassigned nucleic acid, and "\*" indicates a stop codon in the amino acid sequence.

Figure 4A depicts the original overlay of a series of ESTs with some nucleic acid sequence homology to ESTs encoding human placental bikurun, or portions thereof. Shown for reference are the relative positions of bikurun (7-64) and bikurun (102-159), labeled KID1 and KID2 respectively.

Figure 4B depicts a subsequent more comprehensive EST overlay incorporating additional ESTs. Numbers on the upper X-axis refer to length in base pairs, starting at the first base from the most 5' EST sequence. The length of each bar is in proportion to the length in base pairs of the individual ESTs including gaps. The EST accession numbers are indicated to the right of their respective EST bars.

Figure 4C depicts the corresponding alignment of the oligonucleotide sequences of each of the overlapping ESTs shown schematically in Figure 4B. The upper sequence (SEQ ID NO: 51) labeled bikunin represents the consensus

and 1005) are base insertions observed in R74593 that were consistently absent in each of the other overlapping ESTs.

Figure 4D term is the amore and translation of the

oligonucleotide sequence for bikunin depicted in Figure 4C (SEQ ID NO: 45)

Figure 4E depicts the nucleotide sequence (SEQ ID NO: 46) and corresponding amino acid translation (SEQ ID NO: 47) of a placental bikurun encoding sequence that was derived from a human placental cDNA library by PCR-based amplification.

Figure 4F depicts the nucleotide sequence (SEQ ID NO: 48) and corresponding amino acid translation (SEQ ID NO: 49) of a native human placental bikunin encoding clone that was isolated from a human placental lambda cDNA library by colony hybridization.

Figure 4G compares the alignment of the amino acid translated oligonucleotide sequences for placental bikunin obtained by EST overlay (SEQ ID NO: 45), PCR based cloning (SEQ ID NO: 47), and conventional lambda colony hybridization (SEQ ID NO: 49).

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Figure 5 shows a graph of purification of human placental bikurun from placental tissue after Superdex 75 Gel-Filtration. The plot is an overlay of the protein elution profile as measured by OD 280 nm (solid line), activity of eluted protein in a trypsin inhibition assay (% inhibition shown by circles), and activity of eluted protein in a kallikrein inhibition assay (% inhibition shown by squares).

Figure 6 shows a graph which plots the purification of human placental bikunin from placental tissue using C18 Reverse-Phase Chromatography. The plot is an overlay of the protein elution profile as measured by OD 215 nm (solid line), activity of eluted protein in a trypsin inhibition assay (% inhibition shown by circles), and activity of eluted protein in a kallikrein inhibition assay (% inhibition shown by squares).

Figure 7 depicts a silver stained SDS-PAGE gel of highly purified placental bikunin (lane 2), and a series of molecular size marker proteins (lane 1) of the indicated sizes in kilodaltons. Migration was from top to bottom.

Figure 8 shows the amount of trypsin inhibitory activity present in the cell-free fermentation broth from the growth of yeast strains SC101 (panel 8A) or WHL341 (panel 8B) that were stably transformed with a plasmid (pS604) that directs the expression of placental bikunin (102-159).

Figure 9 shows both a silver stained SDS-PAGE (left panel) and a Western blot with anti-placental bikunin (102-159) pAb (right panel) of cell-free fermentation broth from the growth of yeast strain SC101 (recombinants 2.4 and 2.5) that was stably transformed with a plasmid directing the expression of either bovine aprotinin, or placental bikunin (102-159). Migration was from top

to bottom.

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Figure 10 is a photograph which shows a silver stained SDS-PAGE of highly purified placental bikurun (102-159) (lane 2) and a series of molecular size marker proteins (lane 1) of the indicated sizes in Kilodaltons. Migration was from top to bottom.

Figure 11 is a photograph which shows the results of Northern blots of mRNA from various human tissues that was hybridized to a <sup>32</sup>P labeled cDNA probe encoding either placental bikunin (102-159) (panel 11A) or encoding placental bikunin (1-213) (panel 11B). Migration was from top to bottom. The numbers to the right of each blot refer to the size in kilobases of the adjacent RNA markers. The organs from which mRNA was derived is described under each lane of the blot.

Figure 12 depicts an immunoblot of placental derived placental bikunin with rabbit antiserum raised against either synthetic reduced placental bikunin (7-64) (panel A) or 102-159 (panel B). For each panel, contents were: molecular size markers (lanes 1); native placental bikunin isolated from human placenta (lanes 2); synthetic placental bikunin (7-64) (lanes 3) and synthetic placental bikunin (102-159) (lanes 4). Tricine 10-20% SDS-PAGE gels were blotted and developed with protein A-purified primary polyclonal antibody (8 ug IgG in 20 ml 0.1% BSA/Tris-buffered saline (pH 7.5), followed by alkaline phosphatase-conjugated goat anti-rabbit secondary antibody. Migration was from top to bottom.

Figure 13 depicts a Coomassie Blue stained 10-20% Tricine SDS-PAGE gel of 3 micrograms of highly purified placental bikunin (1-170) derived from a baculovirus / Sf9 expression system (lane 2). Lane 1 contains molecular size markers. Migration was from top to bottom.

Figure 14 depicts a comparison of the effect of increasing concentrations of either Sf9-derived human placental bikunin (1-170) (filled circles), synthetic placental bikunin (102-159) (open circles), or aprotinin (open squares) on the activated partial thromboplastin time of human plasma. Clotting was initiated with CaCl<sub>2</sub>. The concentration of proteins are plotted versus the -fold

#### Petaned Description of the invention

The present invention encompasses a new widentified human protein herein called human placental bikunin that contains two serine protease which is tomained the Kunitz lass. The metal refer has a serine protease.

pharmaceutical compositions containing placental bikunin and fragments thereof that are useful for the reduction of perioperative blood loss in a patient undergoing surgery, or with major trauma.

The present invention also provides methods for reducing perioperative blood loss in a patient undergoing surgery or due to major trauma, wherein an effective amount of the disclosed human serine protease inhibitors of the present invention, in a biologically compatible vehicle, is administered to the patient.

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A preferred application for placental bikunin, isolated domains, and other variants is for the reduction of blood loss resulting from trauma or surgery that has the potential for loss of large volumes of blood. These methods and compositions reduce or eliminate the need for whole donor blood or blood products, thereby reducing the risk of infection and other adverse side effects, as well as the cost of surgery. The methods are thus useful in reducing blood loss in normal patients, i.e., those not suffering from inborn or other preoperative deficiencies in coagulation factors. The reduction in blood loss is seen as a reduction in blood loss during surgery, as reduced post surgical drainage or both. Preferred surgical applications include but are not limited to use in thoracic and abdominal surgery, total and partial hip replacement surgeries and surgeries to treat a patient having an epithelial lesion of the eye. Preferred thoracic surgical procedures include but are not limited to aortocoronary bypass, excision of cardiac and aortic aneurysms, and surgery for esophageal varices, and coronary artery bypass surgery. Preferred abdominal surgeries include but are not limited to liver transplants, radical prostatectomy, surgery for diverticulitis of colon, tumor debulking, surgery on the abdominal aorta and surgery for duodenal ulcers, and repair of liver or spleen trauma. Preferred use for the treatment of trauma include but are not limited to the use in stabilization of severely injured patients at accident sites suffering from e.g., limb loss or major thoracic /abdominal wounds. In case of use for the reduction of blood loss resulting from surgery it is preferred to administer the placental bikunin, isolated domains, or other variant prior to and during surgery, whereas in case of use in trauma settings the placental bikurun variant, isolated domain or other variant is to be administered as rapidly as possible following injury, and should be contained on emergency vehicles traveling to the accident sites.

Factor XII (also known as Hageman Factor) is a serine protease that is found in the circulation in a zymogen form (80 kD) at approximately 29-40

μg/ml (see Pixley, et al. (1993) Meth. in Enz., 222, 51-64) and is activated by tissue and plasma kallikrein. Once activated, it participates in the intrinsic pathway of blood coagulation which is activated when blood or plasma contacts a "foreign" or anionic surface. Once activated, Factor XIIa can then cleave and activate a number of other plasma proteases including Factor XI, prekallikrein, and C1 of the complement system. Thus Factor XII may be involved in causing hypotensive reactions since activated kallikrein can cleave kininogen releasing bradykinin (see Colman, (1984) J. Clin. Invest., 73, 1249).

Sepsis is a disease that results from bacterial infection due to bacterial endotoxin or lipopolysaccharide (LPS). Exposure of Factor XII to LPS results in the activation of Factor XII. Patients with sepsis frequently have symptoms of intravascular coagulation which may also be due to activation of Factor XII by LPS. Septic shock can result from bacterial infection and is associated with fever, low systemic vascular resistance, and low arterial pressure. It is a common cause of death in intensive care units in the United States, where seventy five percent of the patients that die from septic shock have a persistent hypotension (see Parillo, et al. (1989) *Ann Rev. Med.*, 40, 469-485).

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Adult respiratory distress syndrome is characterized by pulmonary edema, hypoxemia, and decreased pulmonary compliance. The pathogenesis of the disease is currently unknown although the proteolytic pathways of coagulation and fibrinolysis are believed to play a role (see Carvalho, et al. (1988) J. Lab Clin. Med., 112: 270-277).

The proteins of the instant invention are also a novel human Kunitz type inhibitor of kallikrein, an activator of Factor XII. Thus another object of the current invention is to present a method for the prophylactic or therapeutic treatment of systemic inflammatory reactions such as septic shock, adult respiratory distress syndrome (ARDS), preeclampsia, multiple organ failure and disseminated intravascular coagulation (DIC). The therapeutic or prophylactic administration of the peptides of the instant invention would result in the modulation of these inflammatory conditions and be beneficial to the patient.

professed mediate migration of and tissue invasion by both endotheral considering angiogenesis/neovascularization, and cancer cells during metastasis. Neovascularization is essential to support tumor growth and metastasis is a professed when mediates for spreading of tumors and wring a passecular continuous.

extremely poor patient prognosis.

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Several preclinical studies suggest that Kunitz like serine protease inhibitors with a protease specificity similar to aprotinin are useful as medicaments for cancer. For example, aprotinin reduced tumor growth and invasion, with increased tumor necrosis when administered to hamsters bearing a highly invasive fibrosarcoma or to mice bearing a similarly malignant mammary carcinoma (Latner et al., (1974), Br. J. Cancer 30: 60-67; Latner and Turner, (1976), Br. J. Cancer 33: 535-538). Furthermore, administration of 200,000 KIU of aprotinin i.p. to C57B1/6 Cr male mice on days 1 to 14 postinoculation with Lewis lung carcinoma cells, reduced pulmonary metastases by 50% although had no effect on primary tumor mass (Giraldi et al., (1977) Eur. J. Cancer, 13: 1321-1323). Similarly, administration of 10,000 KIU i.p. on each of days 13-16 post-inoculation of C57BL/6J mice with Lewis tumor cells inhibited pulmonary metastases by 90% without affecting the primary tumor growth (Uetsuji et al., (1992), Jpn. J. Surg. 22: 429-442). In this same study, administration of plasmin or kallikrein with the same dosing schedule was argued to increase the number of pulmonary metastases. These results prompted the authors to suggest that perioperative administration of aprotinin to cancer patients may reduce the likelihood of metastases. Black and Steger (1976, Eur. J. Pharmacol., 38: 313-319) found that aprotinin inhibited the growth of the transplanted rodent Murphy-Strum lymphosarcoma in rats and suggested that the effect involved the inhibition of the kinin-forming enzyme system. Twice daily i.p. injection of female ddY mice with 10,000 KIU of aprotinin for 7 weeks to mice each bearing a single autochtonous squamous cell carcinoma resulting from 3-methylcholanthrene treatment reduced the growth rate of the primary tumors by 90%. In some animals tumor regression was observed. While all vehicle treated animals had died within the seven weeks, all of the aprotinin treatment group remained alive. Reduced tumor growth was associated with hyperkeratosis (Ohkoshi, Gann (1980), 71: 246-250).

Clinically, a surgically cured group of 26 patients who received aprotinin i.v. exhibited a 70% survival two years post surgery with no recurrence of tumors whereas a placebo group of 26 patients at the same time exhibited only a 38% survival with a significant rate of tumor recurrence (Freeman et al. Br. Soc. Gastroenterol. (1980) supplement A: 902). In a case study (Guthrie et al., Br. J. Clin. Pract (1981) 35: 330-332), administration of bromocriptine plus aprotinin to a patient with advanced cancer of the cervix caused remission. Aprotinin was administerd both as a 500,000 KIU i.p. bolus every eight hours concurrently

with a continuous i.v. infusion of aprotinin at a rate of 200,000 KIU per 6 hr for a total of seven days once a month. Treatment was ended at the end of the fourth month due to the development of an allergic reaction to aprotinin. More recent evidence has further underscored a role of plasmin as a target for these effects of aprotinin on metastases.

The mechanism for these events could be related to the fact that aprotinin blocks the invasive potential of cancer cell lines (Liu G., et al., Int J. Cancer (1995), 60: 501-506). Furthermore, since the proteins of the instant invention are also potent inhibitors of plasmin and kallikrien, they are contemplated for use as anti-cancer agents. For example they are contemplated for use in blocking primary tumor growth by restricting neovascularization, primary tumor invasion and in blocking metastasis through inhibition of tissue infiltration. The compounds may be administered locally to tumors or systemically. In a preferred mode of treatment, the protein would be administered perioperatively during tumor debulking to minimize the risk of metastasis. In such a regime, the blood sparing properties of the compound would be additionally advantageous in providing a clearer surgical field of view. Another preferred mode of administration would be as a combination therapy with either MMP inhibitors or chemotherapy. An additional preferred mode of administration would be as a locally administered gene therapy designed to achieve selective expression of placental bikunin within the tumor cells, or their associated stroma and vascular beds.

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Preferred types of cancers targeted for therapy would be vasular-dependent solid tumors such as breast, colon, lung, prostate and ovarian carcinomas which exhibit a high metastatic potential, and those for which local delivery of a high concentration of the protein is feasible such as lung cancers through pulmonary delivery, colon carcinomas through hepatic delivery to liver metastasis, or skin cancers such as head and neck carcinomas or melanomas through subcutaneous delivery. Since the proteins of the present invention are of human origin they would be less likely to be associated with allergic or anaphylactic reactions of the kind observed by Guthrie et al., supra-

activation of the intrinsic pathway of coagulation. This would include prevention of pulmonary embolism in late stage cancer patients a frequent task in team of part. Alb 1964. Haenowiss 24 128-131

Edema of the brain and spinal cord is a complication resulting from traumatic brain or spinal cord injury, stroke, cerebral ischemia, cerebral and sub-arachnoid hemhorrhage, surgery (including open heart surgery), infectious diseases such as encephalitis and meningitis, granulomatous diseases such as Sarcoid and focal or diffuse carcinomas, and is a contributor to the high level of morbidity and death following these events. Bradykinin is known to disrupt the blood brain barrier experimentally (Greenwood J., (1991), Neuroradiology, 33: 95-100; Whittle et al., (1992), Acta Neurochir., 115: 53-59), and infusion of bradykinin into the internal carotid artery induced brain edema in spontaneously hypertensive rats (SHR) subjected to common carotid artery occlusion (Kamiya, (1990), Nippon Ika Daigaku Zasshi. 57: 180-191). Elevated levels of bradykinin are found in extracellular fluids following trauma in a model involving traumatized rat spinal chord (Xu et al., (1991), J. Neurochem, 57: 975-980), and in plasma and tissue from rats with brain edema resulting from cerebral ischaemia (Kamiya et al., (1993), Stroke, 24: 571-575). Bradykinin is released from high molecular weight kininggen by serine proteases including kallikrein (Coleman (1984) J. Clin Invest., 73: 1249), and the serine protease inhibitor aprotinin was found to block the magnitude of brain edema resulting from cerebralschemia in SHR rats (Kamiya, (1990), Nippon Ika Daigaku Zasshi. 57: 180-191; Kamiya et al., (1993), Stroke, 24: 571-575) and rabbits subjected to a cold lesion of the brain (Unterberg et al., (1986), J. Neurosurgery, 64: 269-276).

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These observations indicate that brain edema results from local proteolytic release of kinins such as bradykinin from high molecular weight kininogen, followed by bradykinin-induced increases in blood brain barrier permeability. Accordingly, placental bikunin and fragments thereof are contemplated as medicaments for the prevention of edema in patients at risk for this condition, particularly those of high risk of mortality or brain injury. This would include head and spinal trauma patients, polytrauma patients, patients undergoing surgery of the brain or spinal cord and their associated vessels or other general surgeries including open-heart surgery, patients who have suffered from a stroke, cerebral or sub-arachnoid hemorrhage, infectious diseases of the brain, granulomatous disease of the brain or diffuse or focal carcinomas and tumors of the brain or any conditions such as multiple sclerosis involving breakdown of the blood brain barrier or patients suffering from any other inflammatory processes of the brain or spinal cord. Patients would receive an administration of placental bikunin either as an infusion or bolus injection, intravenously or intracranially. Additional doses of placental bikunin

could be administered intermittently over the following one to three weeks. Dose levels would be designed to attain circulating concentrations in excess of those required to neutralize elevations in plasma levels or bradykinin and other vasoactive peptides formed through the action of serine proteases, and sufficient to reduce edema. Since the protein is of human origin, repeated administration in this course of therapy would not lead to development of an immune reaction to the protein. Placental bikunin and fragments thereof would be contemplated for monotherapy or prophylacsis as well as for use in combination with other medicaments such as neurotherapeutics and neuroprotectants.

Recent evidence (Dela Cadena R. A. at al., (1995), FASEB J. 9: 446-452) has indicated that the contact activation pathway may contribute to the pathogenesis of arthritis and anemia, and that kallikrein inhibitors may be of therapeutic benefit. Accordingly, protease inhibitors of the present invention are contemplated according to their capacity to inhibit human kallikrein, as medicaments for the treatment of arthritis and anemia in humans.

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Treatment of male non-insulin diabetic (NIDDM) patients with aprotinin significantly improved total glucose uptake and decreased the metabolic clearance rate of insulin (Laurenti et al., (1996), Diabetic Medicine 13: 642-645). Accordingly, the human proteins of the present invention are contemplated for chronic use as medicaments for the treatment of NIDDM.

Daily treatment of patients at risk of preterm delivery with urinary trypsin inhibitor for two weeks significantly reduced recurrent uterine contractions (Kanayama et al., (1996), Eur J. Obstet Gynecol. & Reprod Biol 57 133-138). Accordingly, the human proteins of the present invention are contemplated for use in the prevention of preterm delivery

Aprotinin has been shown to stimulate differentiation of mouse myoblasts in culture (Wells and Strickland, Development, (1994), 120: 3639-3647)), a process that is inhibited by TGFb. TGFb exists as an inactive propolypeptide which is activated by limited proteolysis. The mechanism of aprotinin action has been proposed to involve inhibition of proteases which

example, TGF-b concentrations paralleled the extent of bleomycin-induced inflammation. Furthermore, plasmin levels in the alreedar main chage and otherwise field account the alreedar main chage.

a-2-antiplasmin abrogated the post translational activation of pro-TGFb by the macrophage (Khal et al., (1996), Am. J. Respir. Cell Mol. Biol. 15: 252-259.) The data suggest that plasmin contributes to the formation of active TGFb by alveolar macrophage, and that this process plays a pathologic role in the bleomycin-induced lung inflammation.

In light of these observations, placental bikunin and fragments thereof are contemplated as therapeutics for various fibrotic disorders, including pulmonary, hepatic, renal and dermal (scleroderma) fibrosis.

Aerosilized aprotinin was shown to protect >50% of mice infected with lethal doses of either influenza virus or paramyxovirus (Ovcharenko and Zhirnov, Antiviral Research, (1994), 23: 107-118). A suppression of the development of fatal hemorrhagic bronchopneumonia and a normalization of body weight gain were also noted with aerosilized aprotinin treatment. In light of these observations, placental bikunin and fragments thereof are contemplated as therapeutics for various respiratory related influenza-like diseases.

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The human placental bikunin, isolated domains, and other variants of the invention are contemplated for use in the medical/therapeutic applications suggested for native aprotinin or aprotinin analogues with other inhibitory profiles, in particular those which necessitate usage of large doses. These would include diseases for which use of the human protein is indicated by virtue of its ability to inhibit human serine proteases such as trypsin, plasmin, kallikrein, elastase, cathepsin G and proteinase-3, which include and are not limited to: acute pancreatitis (pancreatic elastase and trypsin), inflammation, thrombocytopenia, preservation of platelet function, organ preservation, wound healing, various forms of shock, including shock lung, endotoxin shock and post operative complications; disturbances of blood coagulation such as hyperfibrinolytic hemorrhage; acute and chronic inflammatory reactions, in particular for the therapy and prophylaxis of organ lesions, such as for example pancreatitis and radiation induced enteritis, complex-mediated inflammatory reactions such as immunovasculitis, glomerulonephritis and types of arthritis; collagenoses in particular rheumatoid arthritis; types of arthritis caused by metabolism-related deposits (for example gout); degeneration of the elastic constituents of the connective tissue parts of organs, such as in atherosclerosis (serum elastase) or pulmonary emphysema (neutrophil elastase); adult respiratory distress syndrome, inflammatory bowel disease, and psoriasis.

A major unexpected finding was that the synthetic peptides encoding

bikurun (7-64), and bikurun (102-159), could properly fold into the correct three-dimensional conformation having active protease inhibitor bioactivity (Examples 2 and 1, respectively). Upon folding, each of these fragments of Bikurun underwent a reduction in mass of 6 mass units, consistent with the formation in each case, of three intrachain disulfide bonds between six cysteine residues of each fragment. Another surprising finding is that the synthetic peptides encoding bikurun (7-64), bikurun (102-159), and bikurun (1-170) are highly inhibitory of plasmin and both tissue and plasma kallikrein (Example 4, 3, and 10 respectively). Inhibition of plasmin and kallikrein by Trasylol® is thought to be involved in the mechanism by which Trasylol® reduces blood loss during open heart surgery. Our unexpected findings of the specificity of the Kunitz domains of the present invention make them suitable therapeutic agents for blood sparing during surgery or trauma where there is significant blood loss, or for any other condition where inhibition of plasmin and/or kallikrein would be beneficial.

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Furthermore, we showed in this disclosure (Example 10) that placental bikunin (1-170) is a potent inhibitor of factor XIa and a moderate inhibitor of factor Xa. Factor XIa plays an essential role in the intrinsic pathway of coagulation, serving to interconvert inactive factor IX into active factor IXa. Thus, Placental Bikunin inhibits two key enzymes of the intrinsic pathway, kallikrein and factor XIa. Consistent with these observations, we also showed that placental bikunin (1-170) is a potent inhibitor of the activated partial thromboplastin time, which is a measure of the speed of coagulation driven by the intrinsic pathway. On the other hand, we showed that Placental bikunin (1-170) is an extremely weak inhibitor of the tissue factor VIIa complex, suggesting that it is not important in the regulation of the extrinsic coagulation cascade Based on these unexpected findings, placental bikunin is contemplated as a medicament for diseases in which activation of the intrinsic pathway of coagulation contributes significantly to the disease mechanism. Examples of such diseases would include post-traumatic shock and disseminated intravascular coagulation.

large doses of the proteins. Being of human origin, the protein of the instant invention can thus be administered to human patients with significantly and a contract of the instant origin.

administration of similar doses of Trasylol<sup>®</sup> Furthermore, it was found that bikunin (102-159), bikunin (7-64), and bikunin (1-170) are significantly more potent inhibitors of plasma kallikrein than Trasylol<sup>®</sup> in vitro (Example 3, 4 and 10). Thus bikunin and fragments thereof are expected to be more effective in vitro at lowering blood loss in patients.

The amount of serine protease inhibitor administered should be sufficient to provide a supra normal plasma level. For the prophylactic reduction of bleeding during and following coronary aortic by-pass surgery (CABG), the proteins of the instant invention may be used in place of Trasylol® while taking into account the differences in potency. The use of Trasvlol® is outlined in the Physicians Desk Reference, (1995), listing for Trasylol® supplement A. Briefly, with the patient in a supine position, the loading dose of placental bikunin, isolated domain or other variant is given slowly over about 20 to 30 minutes, after induction of anesthesia but prior to sternotomy. In general, a total dose of between about 2x106 KIU (kallikrein inhibitory units) and 8 X106 KIU will be used, depending on such factors as patient weight and the length of the surgery. Preferred loading doses are those that contain a total of 1 to 2 million kallikrein inhibitory units (KIU). When the loading dose is complete, it is followed by the constant infusion dose, which is continued until surgery is complete and the patient leaves the operating room. Preferred constant infusion doses are in the range of about 250,000 to 500,000 KIU per The pump prime dose is added to the priming fluid of the cardiopulmonary bypass circuit, by replacement of an aliquot of the priming fluid prior to the institution of the cardiopulmonary bypass. Preferred pump prime doses are those that contain a total of about one to two million KIU

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The proteins of the instant invention are employed in pharmaceutical compositions formulated in the manner known to the art. Such compositions contain active ingredient(s) plus one or more pharmaceutically acceptable carriers, diluents, fillers, binders, and other excipients, depending on the administration mode and dosage form contemplated. Examples of therapeutically inert inorganic or organic carriers known to those skilled in the art include, but are not limited to, lactose, corn starch or derivatives thereof, talc, vegetable oils, waxes, fats, polyols such as polyethylene glycol, water, saccharose, alcohols, glycerin and the like. Various preservatives, emulsifiers, dispersants, flavorants, wetting agents, antioxidants, sweeteners, colorants, stabilizers, salts, buffers and the like can also be added, as required to assist in the stabilization of the formulation or to assist in increasing bioavailability of

the active ingredient(s) or to yield a formulation of acceptable flavor or odor in the case of oral dosing. The inhibitor employed in such compositions may be in the form of the original compound itself, or optionally, in the form of a pharmaceutically acceptable salt. The proteins of the instant invention can be adminstered alone, or in various combinations, and in combination with other therapeutic compositions. The compositions so formulated are selected as needed for administration of the inhibitor by any suitable mode known to those skilled in the art.

Parenteral administration modes include intravenous (i.v.), subcutaneous (s.c.), intraperitoneal (i.p.), and intramuscular (i.m.) routes. Intravenous administration can be used to obtain acute regulation of peak plasma concentrations of the drug as might be needed. Alternatively, the drug can be administered at a desired rate continuously by i.v. catheter. Suitable vehicles include sterile, non-pyrogenic aqueous diluents, such as sterile water for injection, sterile-buffered solutions or sterile saline. The resulting composition is administered to the patient prior to and/or during surgery by intravenous injection or infusion.

Improved half-life and targeting of the drug to phagosomes such as neutrophils and macrophage involved in inflammation may be aided by entrapment of the drug in liposomes. It should be possible to improve the selectivity of liposomal targeting by incorporating into the outside of the liposomes ligands that bind to macromolecules specific to target organs / tissues such as the GI tract and lungs. Alternatively, i.m. or s.c. deposit injection with or without encapsulation of the drug into degradable microspheres (e.g., comprising poly-DL-lactide-co-glycolide) or protective formulations containing collagen can be used to obtain prolonged sustained drug release. For improved convenience of the dosage form it is possible to use an in implanted reservoir and septum such as the percuseal system. Improved convenience and patient compliance may also be achieved by use of either injector pens (e.g., the Novo Pin or Q-pen) or needle-free jet injectors (e.g., from Bioject, Mediject or Becton Dickinson). Precisely controlled release can also be achieved using implantable

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Nasal delivery may be achieved by incorporating the drug into bloadhesive particulate carriers (<200 mm) such as those comprising to \$100 km. The such as those comprising to \$100 km.

enhancers such as phospholipids or acylcarnitines. Commercially available systems include those developed by Dan Biosys and Scios Nova.

Pulmonary delivery represents a nonparenteral mode of administration of the drug to the circulation. The lower airway epithelia are highly permeable to a wide range of proteins of molecular sizes up to about 20 kDa. Micron-sized dry powders containing the medicament in a suitable carrier such as manutol, sucrose or lactose may be delivered to the distal alveolar surface using dry powder inhalers such as those of Inhale<sup>TM</sup>, Dura<sup>TM</sup>, Fisons (Spinhaler<sup>TM</sup>), and Glaxo (Rotahaler<sup>TM</sup>), or Astra (Turbohaler<sup>TM</sup>) propellant based metered dose inhalers. Solution formulations with or without liposomes may be delivered using ultrasonic nebulizers.

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Oral delivery may be achieved by incorporating the drug into tablets, coated tablets, dragées, hard and soft gelatin capsules, solutions, emulsions, suspensions or enteric coated capsules designed to release the drug into the colon where digestive protease activity is low. Examples of the latter include the OROS-CT/Osmet<sup>TM</sup> system of ALZA, and the PULSINCAP<sup>TM</sup> system of Scherer Drug Delivery Systems. Other systems use azo-crosslinked polymers that are degraded by colon-specific bacterial azoreductases, or pH sensitive polyacrylate polymers that are activated by the rise in pH in the colon. The above systems may be used in conjunction with a wide range of available absorption enhancers. Rectal delivery may be achieved by incorporating the drug into suppositories.

In its preferred medicinal application, for reduction of perioperative blood loss, the preferred mode of administration of the placental bikunin variants of the present invention is parenterally, preferably by t.v. route through a central line.

The amount of the pharmaceutical composition to be employed will depend on the recipient and the condition being treated. The requisite amount may be determined without undue experimentation by protocols known to those skilled in the art. Alternatively, the requisite amount may be calculated, based on a determination of the amount of target protease such as plasmin or kallikrein which must be inhibited in order to treat the condition. As the active materials contemplated in this invention are deemed to be nontoxic, treatment preferably involves administration of an excess of the optimally required amount of active agent.

Additionally, placental bikunin, isolated domains or other variants may be used to isolate natural substances such as its cognate proteases from human

material using affinity based separation methods, as well as to elicit antibodies to the protease that can be further used to explore the tissue distribution and useful functions of Placental bikunin.

### 5 Searching Human Sequence Data

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The existence of a distinct human protein homologous in function to aprotinin, was deduced following a unique analysis of sequence entries to the expressed-sequence-tag data-base (hereafter termed dbEST) at the NCBI (National Center for Biological Information, Maryland). Using the TBlastN algorithm (BLAST, or Basic Local Alignment Search Tool uses the method of Altschul et a., (1990) J. Mol Biol 215: 403-410, to search for similarities between a query sequence and all the sequences in a data-base, protein or nucleic acid in any combination), the data-base was examined for nucleotide sequences bearing homology to the sequence of bovine pre-pro-aprotinin, Trasylol®. This search of numerous clones was selectively narrowed to two particular clones which could possibly encode for a deduced amino acid sequence that would correspond to a human protein homologous in function to aprotinin. The selected nucleic acid sequences were R35464 (SEQ ID NO: 12) and R74593 (SEQ ID NO: 14) that were generated from a human placental nucleic acid library. The translated protein sequence in the longest open reading frame for R35464 (SEQ ID NO: 13) was missing one of the 6 cysteines that are critical for formation of the Kunitz-domain covalent structure, meaning that the nucleic acid sequence of R35464 could not yield a functional inhibitor. Similarly, the longest translated open reading frame from clone R74593 (SEQ ID NO(15)contained a stop codon 5' to the region encoding the Kunitz like sequence, meaning that this sequence, could not be translated to yield a functional secreted Kunitz domain. The significance of these sequences alone was unclear. It was possible that they represented a) the products of pseudogenes, b) regions of untranslated mRNA, or c) the products of viable mRNA which had been sequenced incorrectly

<sup>35 3&#</sup>x27; to the segment of cDNA encoding our proposed Kurutz like sequences found within R35464 and R74593. The primers used to amplify a fragment one ding one knows like regions of R74593 were.

CGAAGCTTCATCTCCGAAGCTCCAGACG (the 3'primer with a HindIII site, SEQ ID NO:33) and AGGATCTAGACAATAATTACCTGACCAAGGA (the 5'primer with an XbaI site; SEQ ID NO:34).

These primers were used to amplify by PCR (30 cycles) a 500 base pair product from a human placental cDNA library from Clontech (MATCHMAKER, Cat #HL4003AB, Clontech Laboratories, Palo Alto, CA), which was subcloned into Bluescript-SK+ and sequenced with the T3 primer with a Sequenase<sup>TM</sup> kit version 2.0. Surprisingly, the sequence of the fragment obtained using our primers was different from the sequence listed in the dbEST data base for clone R74593. In particular, our new sequence contained an additional guanosine base inserted 3' to the putative stop codon, but 5' to the segment encoding the Kunitz-like sequence (Figure 3). The insertion of an additional G shifted the stop codon out of the reading frame for the Kunitz-like domain (G at base pair 114 of the corrected sequence for R74593; Figure 3).

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Subsequent query of the dbEST for sequences homologous to the Kunitz-like peptide sequence of R74593 yielded H94519 derived from human retina library and N39798. These sequences contained a Kunitz-like sequence that was almost identical to the Kunitz-like domain encoded in R35464 except that it contained all six of the characteristic cysteines. Overlay of each of the nucleotide sequences with that of R74593 (corrected by the insertion of G at b,p, 114) and R35464 was used to obtain a consensus nucleotide sequence for a partial human placental bikunin (SEQ ID NO: 9; Figure 3). The translated consensus sequence yielded an open reading frame extending from residue -18 to +179 (Figure 3; full translation SEQ ID NO: 10) that contained two complete Kunitz-like domain sequences, within the region of amino acid residues 17-64 and 102-159 respectively.

Further efforts attempted to obtain additional 5' sequence by querying dbEST with the sequence of R35464. Possible matches from such searches, that possessed additional 5' sequence were then in turn used to re-query the dbEST. In such an iterative fashion, a series of overlapping 5' sequences were identified which included clones H16866, T66058, R34808, R87894, N40851 and N39876 (Figure 4). Alignment of some of these sequences suggested the presence of a 5' ATG which might serve as a start site for synthesis of the consensus translated protein sequence. From this selected information, it was now possible to selectively screen for, and determine the nucleic acid and polypeptide sequences of a human protein with homologous function to aprotinin.

Re-interrogation of the dbEST revealed a number of new EST entries

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shown schematically in Figure 4B. Overlap with these additional ESTs allowed us to construct a much longer consensus oligonucleotide sequence (Figure 4C) that extended both 5' and 3' beyond the original oligonucleotide sequence depicted in Figure 3. In fact, the new sequence of total length 1.6 kilobases extended all the way to the 3' poly-A tail. The increased number of overlapping ESTs at each base-pair position along the sequence improved the level of confidence in certain regions such as the sequence overlapping with the 3' end of EST R74593 (Figure 3). Several overlapping ESTs in this region corroborated two critical base deletions relative to R74593 (located as bold underlined in Figure 4C, map positions 994 and 1005). Translation of the new consensus sequence (Figure 4D) in the bikunin encoding frame yielded a form of placental bikunin that was larger (248 amino acids) than the mature sequence (179 amino acids) encoded from the original consensus (SEQ ID NO: 1), and was terminated by an in-frame stop codon within the oligonucleotide consensus The size increase was due to a frame shift in the 3' coding region resulting from removal of the two base insertions unique to EST R74593. The frame shift moved the stop codon of the original consensus (Figure 3) out of frame enabling read through into a new frame encoding the additional amino acid sequence. The new translation product (Figure 4D) was identical to the original protein consensus sequence (SEQ ID NO: 1) between residues +1 to +175 (encoding the Kunitz domains), but contained a new C-terminal extension exhibiting a putative 24 residue long transmembrane domain (underlined in Figure 4D) followed by a short 31 residue cytoplasmic domain. The precise sequence around the initiator methionine and signal peptide was somewhat tentative due to considerable heterogeneity amongst the overlapping ESTs in this region.

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Analysis of the protein sequence by Geneworks<sup>TM</sup>, highlighted asparagine residues at positions 30 and 67 as consensus sites for putative N-linked glycosylation. Asparagine 30 was not observed during N-terminal sequencing of the full length protein isolated from human placenta, consistent with it being glycosylated

bikunin nucleotide sequence inferred from the analysis of Figure 3, was confirmed as follows. The nucleic acid primer hybridizing 5' to the Knipitz-

sequence in Figure 3).

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### GGTCTAGAGGCCGGGTCGTTTCTCGCCTGGCTGGGA

(a 5' primer derived from R35464 sequence with an XbaI site; SEQ ID NO: 35), and the nucleic acid primer hybridizing 3' to the Kunitz encoding sequence of R74593 (b.p. 680-700 of consensus nucleotide sequence in Figure 3), was used to PCR amplify, from a Clontech human placental library, a fragment of the size (ca. 670 b.p) expected from a cDNA consensus nucleotide sequence encoding the placental bikunin sequence of Figure 3 (Shown schematically in Figure 4A).

Using a 5' primer hybridizing to a sequence in R87894 that is 126 b.p 5' to the putative ATG start site discussed above, (shown schematically in Figure 4A at b.p. 110) plus the same 3' primer to R74593 as used above, it was possible to amplify a fragment from a Clontech human placental library of the expected size (approximately 872 b.p) predicted by EST overlay (Shown schematically in Figure 4).

Sequencing of the 872 b.p. fragment showed it to contain nucleotide segment corresponding to b.p. 110 to 218 of EST R87894 at its 5' end and b.p. 310 to 542 of the consensus sequence for placental bikunin inferred from the EST overlay analysis (of Figure 3), at its 3' end. This 3' nucleotide sequence contained all of the Kunitz-like domain encoded by placental bikunin (102-159).

To obtain a cDNA encoding the entire extracellular region of the protein, the following 5' PCR primer:

CACCTGATCGCGAGACCCC (SEQ ID NO: 36)

designed to hybridize to a sequence within EST R34808 was used with the same 3' primer to EST 74593 to amplify (30 cycles) an approximately 780 base-pair cDNA product from the human placental cDNA libra: This product was gel purified, and cloned into the TA vector (Invitrogen) for DNA sequencing by the dideoxy method (Sanger F., et al., (1977) Proc. Natl. Acad. Sci (USA), 74: 5463-5467) with the following primers:

30 Vector Specific: GATTTAGGTGACACTATAG (SP6) (SEQ ID NO: 37 TAATACGACTCACTATAGGG (T7) (SEQ ID NO: 38 Gene Specific: TTACCTGACCAAGGAGGAGTGC (SEQ ID NO: 39 AATCCGCTGCATTCCTGCTGGTG (SEQ ID NO: 40) CAGTCACTGGGGCCTTGCGGT (SEQ ID NO: 41)

The resulting cDNA sequence is depicted in Figure 4E together with its

translation product. At the nucleotide level, the sequence exhibited only minor differences from the consensus EST sequence (Figure 4D). Translation of the sequence yielded a coding sequence containing an in-frame initiator ATG site, signal peptide and mature placental bikunin sequence and transmembrane domain. The translated sequence of the PCR product was missing the last 12 amino acid residues from the cytoplasmic domain as a consequence of the choice of selection of the 3' primer for PCR amplification. This choice of 3' PCR primer (designed based on the sequence of R74593) was also responsible for the introduction of an artifactual S to F mutation at amino acid position 211 of the translated PCR-derived sequence. The signal peptide deduced from translation of the PCR fragment was somewhat different to that of the EST consensus.

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To obtain a full length placental bikunin cDNA, the PCR derived product (Figure 4E) was gel purified and used to isolate a non-PCR based full length clone representing the bikunin sequence. The PCR derived cDNA sequence was labeled with <sup>32</sup>P-CTP by High Prime (Boehringer Mannheim) and used to probe a placental cDNA Library (Stratagene, Unizap $^{TM}$   $\lambda$  library) using colony hybridization techniques. Approximately 2 X 106 phage plaques underwent 3 rounds of screening and plaque purification. Two clones were deemed full length (~1.5 kilobases) as determined by restriction enzyme analysis and based on comparison with the size of the EST consensus sequence (see above). Sequencing of one of these clone by the dideoxy method yielded the oligonucleotide sequence depicted in Figure 4F. The translation product from this sequence yielded a protein with inframe initiator methionine, signal peptide and mature placental bikunin sequence. The mature placental bikunin sequence was identical to the sequence of the mature protein derived by translation of the EST consensus although the signal peptide sequence lengths and sequences differed. Unlike the PCR derived product, the cDNA derived by colony hybridization contained the entire ectodomain, transmembrane domain, cytoplasmic domain and in-frame stop codon. In fact, the clone extended all the way to the poly-A tail. The initiator methionine was followed by a hydrophobic signal peptide which was identical to the signal peptide encoded in the PCR

human placenta a soluble fragment of placental bikunin which was also an active protease inhibitor (Example 7). Both the natural protein and the time of the first expressed a first end of propagation and section in the springer.

residue at position 30 based on the recoveries of PTH-amino acids during N-terminal sequencing (Examples 7 and 9).

Based on the above observations, it seems that full length placental bikunin has the capacity to exist as a transmembrane protein on the surface of cells as well as a soluble protein. Other transmembrane proteins that contain Kunitz domains are known to undergo proteolytic processing to yield mixtures of soluble and membrane associated forms. These include two forms of the Amyloid Precursor Protein termed APP751 (Esch F., et al., (1990) Science, 248: 1122-1124) and APP 770 (Wang R., et al., (1991), J. Biol Chem, 266: 16960-16964).

Contact activation is a process which is activated by exposure of damaged vascular surfaces to components of the coagulation cascade. Angiogenesis is a process that involves local activation of plasmin at endothelial surfaces. The specificity of placental bikunin and its putative capacity to anchor to cell surfaces, suggest that the physiologic functions of transmembranous placental bikunin may include regulation of contact activation and angiogenesis.

The amino acid sequences for placental bikurin (7-64), bikurin (102-159), and full length placental bikurin (Figure 4F) were searched against the PIR (Vers. 46.0) and PatchX (Vers. 46.0) protein databases as well as the GeneSeq (Vers. 20.0) protein database of patented sequences using the Genetics Computer Group program FastA. Using the Genetics Computer Group program TFastA (Pearson and Lipman, 1988, Proc. Natl. Acad. Sci. USA 85: 2444-2448), these same protein sequences were searched versus the six-frame translations of the GenBank (Vers. 92.0 with updates to 1/26/96) and EMBL (modified Vers. 45.0) nucleotide databases as well as the GeneSeq (Vers. 20.0) nucleotide database of patented sequences. The EST and STS subsets of GenBank and EMBL were not included in this set of searches. The best matches resulting from these searches contained sequences which were only about 50% identical over their full length to the 58-amino acid protein sequence derived from our analysis of clones R74593 and R35464.

#### Isolation of Human Bikunin

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As mentioned above, synthetic peptides corresponding to bikunin (7-64) and bikunin (102-159) as determined from the translated consensus sequence for bikunin (Figure 3), could be refolded (Examples 2 and 1, respectively) to yield active kallikrein inhibitor protein (Example 4 and 3, respectively). We exploited this unexpected property to devise a purification scheme to isolate

native placental bikurun from human tissue.

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Using a purification scheme which employed kallikrein-sepharose affinity chromatography as a first step, highly purified native potent kallikrein inhibitor was isolated. The isolated native human bikunin had an identical N-terminus (sequenced for 50 amino acid residues) as the sequence predicted by the translation of the consensus nucleic acid sequence (Figure 3) amino acid residues +1 to +50 (Example 7). This confirmed for the first time the existence of a novel native kallikrein inhibitor isolated from human placenta.

Known Kunitz-like domains are listed below. Residues believed to be making contact with target proteases are highlighted as of special interest (bold/underlined). These particular residues are named positions Xaa<sup>1-16</sup> for specific reference as shown by label Xaa below:

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Xaa
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                   2 1 456789
      1 IHDFOLUSHUV GRORASMPRW WYNUTDGSOD DFVYGGODGN SMRYLTHEED LYKOATV
      2: YEEYCTANAVT GPORASPPRW YFDVERMSON NFIYGCORGN KMSYRSEEAC MLFOFRQ
      3 - HSFCAFKADD GPOKAIMKRF FFNIFTEQCE EFIYGGCEGN QURFESLEEC KKMOTFD
      4) - PDFCFLEEDP GICRGYITRY FYNNQTKQCE SFKYGGCLGN MARFETLEEC KNICEDG
     5: - PSWCLTPADR GLORANENRE YYNSVIGKOR FERYSGOGGN ENNETSKOEG LRACKEG
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      6: - AEICLLPLDY GPORALLLRY YYRYFTGSGF QFLYGGCEGN AMDFYTWEAG DDAGWFI
      T) -PSFCYSPKDE GLOSANVTRY YFNEFYFTOO AFTYTGOGGN EMMFVSREDO KRACAMA
      8) - KAVOSQEAMT SPORAVMPRT TEDLSYGHOV FEITGGOGGN REDE ESEDYO MAVOHAM
      9 RPDFCLEPPYT GPCKARIIRY FYNAKAGLOQ TFVYGGCRAK RWWKKSAEDC MRTGGGA
    10 - ----CQLGYSA OPOMGMTSRY FYNGTSMACE TEQYGOOMGN SMAFF VTEHEC LOTO
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    11 VAACHURIVR GPORAFIQLW AFDAUFGROU DE PYGGOQCN REFEYED PEYDGUE
    10 - EVOCSEÇAET OPORAMISEW YFOVTESHOA OF FYGGOGGN FRANF DTEEYS MAVOSCA
    13 ----THIPHDE STORDFILKW WYSENTESSA REWYSGOGGN ENFEGSOMES ERWS
    14 - PRINCAFEMEK GPOQTYMTRW FFNFETGECE LFAYGGOGGN SNOWLEHERD EMPOYET
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Where sequence number 1) is Bikunin (7-64) (SEQ ID NO: 4), sequence 2) is Bikunin (102-159) (SEQ ID NO: 6), sequence 3) is Tissue factor pathway

inhibitor precursor 2 (SEQ ID NO 21), sequence 7) is Tissue factor pathway inhibitor precursor 2 (SEQ ID NO 22), sequence 8) is Amyloid precursor to the homologue SEQ ID NO 23, sequence 8) is Amyloid precursor to the homologue SEQ ID NO 23, sequence 8, Appetitus SEQ II No 24.

sequence 10) is Inter- $\alpha$ -trypsin inhibitor precursor (SEQ ID NOs: 25); sequence 11) is Inter- $\alpha$ -trypsin inhibitor precursor (SEQ ID NOs: 26); sequence 12) is Amyloid precursor protein (SEQ ID NO: 27); sequence 13) is Collagen  $\alpha$ -3(VI) precursor (SEQ ID NO: 28); and squence 14) is FiKI-B9 (SEQ ID NO: 29).

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It can be seen that Placental Bikunin (7-64) and (102-159) each have the same number (six) and spacing of cysteine residues as is found in members of the Kunitz class of serine protease inhibitors. The precise bonding of cysteine residues to form the three intrachain disulfide bonds is known and invarient for all previously known Kunitz family members (Laskowski, M et al., 1980, Ann. Rev. Biochem. 49:593-626). Based on this known bonding pattern and the fact that the folding of Placental Bikunin (7-64) and (102-159) into active protease inhibitors is accompanied by a mass reduction consistent with the formation of three intrachain disulfide bonds (Examples 2 and 1), it is highly probable that the disulfide bonding within the Kunitz domains of Placental Bikunin occur between cysteine residues: C11 and C61; C20 and C44; C36 and C57; C106 and C156; C115 and C139; C131 and C152. Furthermore, this pattern of disulfide bonding is highly probable in larger forms of Placental Bikunin containing both Kunitz domains since such forms of the protein are also active serine protease inhibitors and because N-terminal sequencing (Example 7) of native Placental Bikunin for 50 cycles yielded a sequence that was silent at positions where the cysteine residues were expected.

The placental bikunin, isolated domains or other variants of the present invention may be produced by standard solid phase peptide synthesis using either t-Boc chemistry as described by Merrifield R.B. and Barany G., in: The peptides. Analysis, Synthesis, Biology, 2, Gross E. et al., Eds. Academic Press (1980) Chapter 1; or using F-moc chemistry as described by Carpino L.A., and Han G.Y., (1970) J. Amer Chem Soc., 92, 5748-5749, and illustrated in Example 2. Alternatively, expression of a DNA encoding the placental bikunin variant may be used to produce recombinant placental bikunin variants.

The invention also relates to DNA constructs that encode the Placental bikumin protein variants of the present invention. These constructs may be prepared by synthetic methods such as those described in Beaucage S.L. and Caruthers M.H., (1981) Tetrahedron Lett, 22, pp1859-1862; Matteucci M.D and Caruthers M.H., (1981), J. Am. Chem. Soc. 103, p 3185; or from genomic or cDNA which may have been obtained by screening genomic or cDNA libraries with cDNA probes designed to hybridize with placental bikunin encoding DNA sequence. Genomic or cDNA sequence can be modified at one or more

sites to obtain cDNA encoding any of the amino acid substitutions or deletions described in this disclosure.

The instant invention also relates to expression vectors containing the DNA constructs encoding the placental bikunin, isolated domains or other variants of the present invention that can be used for the production of recombinant placental bikunin variants. The cDNA should be connected to a suitable promoter sequence which shows transcriptional activity in the host cell of choice, possess a suitable terminator and a poly-adenylation signal. The cDNA encoding the placental bikunin variant can be fused to a 5' signal peptide that will result in the protein encoded by the cDNA to undergo secretion. The signal peptide can be one that is recognized by the host organism. In the case of a mammalian host cell, the signal peptide can also be the natural signal peptide present in full length placental bikunin. The procedures used to prepare such vectors for expression of placental bikunin variants are well known in the art and are for example described in Sambrook et al., Molecular Cloning: A laboratory Manual, Cold Spring Harbor, New York, (1989).

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The instant invention also relates to transformed cells containing the DNA constructs encoding the placental bikunin, isolated domains or other variants of the present invention that can be used for the production of recombinant placental bikunin variants. A variety of combinations of expression vector and host organism exist which can be used for the production of the placental bikunin variants. Suitable host cells include baculovirus infected Sf9 insect cells, mammalian cells such as BHK, CHO, Hela and C-127, bacteria such as E. coli, and yeasts such as Saccharomyces cervisiae. Methods for the use of mammalian, insect and microbial expressions systems needed to achieve expression of placental bikunin are well known in the art and are described, for example, in Ausubel F M et al., Current Protocols in Molecular Biology, John Wiley & Sons (1995), Chapter 16. For fragments of placental bikunin containing a single Kunitz inhibitor domain such as bikunin (7-64) and (102-159), yeast and E. coli expression systems are preferable, with yeast systems being most preferred. Typically, yeast expression would be carried out

<sup>5,032,573.</sup> Use of mammalian and yeast systems are most preferred for the expression of larger placental bikunin variants containing high profit of tomacon of larger placental bikunin variants.

DNA encoding variants of placental bikurun that possess amino acid substitution of the natural amino sequence can be prepared for expression of recombinant protein using the methods of Kunkel T.A., (1985) Proc. Natl. Acad. Sci USA 82: 488-492. Briefly, the DNA to be mutagenized is cloned into a single stranded bacteriophage vector such as M13. An oligonucleotide spanning the region to be changed and encoding the substitution is hybridized to the single stranded DNA and made double stranded by standard molecular biology techniques. This DNA is then transformed into an appropriate bacterial host and verified by dideoxynucleotide sequencing. The correct DNA is then cloned into the expression plasmid. Alternatively, the target DNA may be mutagenized by standard PCR techniques, sequenced, and inserted into the appropriate expression plasmid.

The following particular examples are offered by way of illustration, and not limitation, of certain aspects and preferred embodiments of the instant invention.

### Example 1

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## Preparation of synthetic placental bikunin (102-159)

Materials and methods/Reagents used. The fluorogenic substrate Tos20 Gly-Pro-Lys-AMC was purchased from Bachem BioScience Inc (King of Prussia, PA). PNGB, Pro-Phe-Arg-AMC, Ala-Ala-Pro-Met-AMC, bovine trypsin (type III), human plasma kallikrein, and human plasmin were from Sigma (St. Louis, MO).

Recombinant aprotinin (Trasylol®) was from Bayer AG (Wuppertal, Germany). Pre-loaded Gln Wang resin was from Novabiochem (La Jolla, CA). Thioanisole, ethanedithiol and t-butyl methyl ether was from Aldrich (Milwaukee, WI).

### Quantification of functional placental bikunin (7-64) and (102-159)

The amount of trypsin inhibitory activity present in the refolded sample at various stages of purification was measured using GPK-AMC as a substrate. Bovine trypsin (200 pmoles) was incubated for 5 min at 37% With bikunin (7-64) or (102-159), from various stages of purification, in buffer A (50 mM Hepes, pH 7.5, 0.1 M NaCl, 2 mM CaCl<sub>2</sub> and 0.01% triton X-100). GPK-AMC was added (20  $\mu$ M final) and the amount of coumarin produced was determined by measuring the fluorescence (ex = 370 nm, em = 432 nm) on a Perkin-Elmer LS-50B fluorimeter over a 2 min. period. For samples being tested the % inhibition

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for each was calculated according to equation 1; where  $R_0$  is the rate of fluorescence increase in the presence of inhibitor and  $R_1$  is the rate determined in the absence of added sample. One unit of activity for the inhibitor is defined as the amount needed to achieve 50% inhibition in the assay using the conditions as described.

% inhibition = 
$$100 \times [1 - R_0/R_1]$$
 (1)

Biosystems model 420A peptide synthesizer using NMP-HBTU Fmoc chemistry. The peptide was synthesized on pre loaded Gln resin with an 8-fold excess of amino acid for each coupling. Cleavage and deprotection was performed in 84.6% trifluoroacetic acid (TFA), 4.4% thioanisole, 2.2% ethanedithiol, 4.4% liquified phenol, and 4.4% H2O for 2 hours at room temperature. The crude peptide was precipitated, centrifuged and washed twice in t-butyl methyl ether. The peptide was purified on a Dynamax 60A C18 reverse-phase HPLC column using a TFA/acetonitrile gradient. The final preparation (61.0 mg) yielded the correct amino acid composition and molecular mass by Electrospray mass spectroscopy (MH+ =6836.1; calcd = 6835.5) for the predicted sequence:

YEEYCTANAV TGPCRASFPR WYFDVERNSC NNFIYGGORG NKNSYRSEEA OMLROFRQ SEQ ID NO: 6)

Purification. Refolding of placental bikunin (132-159) was performed according to the method of Tam et al., (J. Am. Chem. Soc. 1991, 113: 5657-62). A portion of the purified peptide (15.2 mg) was dissolved in 4.0 ml of 0.1 M Tris. pH 6.0, and 8 M urea. Oxidation of the disulfides was accomplished by dropwise addition of a solution containing 23% DMSO, and 0.1 M Tris. pH 6.0 to obtain a final concentration of 0.5 mg/ml peptide in 20% DMSO, 0.1 M Tris. pH 6.0, and 1 M urea. The solution was allowed to stir for 24 hr at 25°C after which it was diluted 1:10 in buffer containing 50 mM Tris. pH 8.0, and 0.1 M

instructions. The refolded material was loaded onto the affinity column at a flow rate of 1 ml imin and washed with 50 mM Tris. pH 50 and 0.1 M National programme at 250 pm or the washed with 10 mg/strip.

column was eluted with 3 volumes each of 0.2 M acetic acid, pH 40 and 1.7 Active fractions were pooled (see below) and the pH of the solution adjusted to 2.5. The material was directly applied to a Vydac C18 reverse-phase column (5 micron, 0.46 x 25 cm) which had been equilibrated in 22.5% acetonitrile in 0.1% TFA. Separation was achieved using a linear gradient of 22.5 to 40% acetonitrile in 0.1% TFA at 1.0 ml/min over 40 min. Active fractions were pooled, lyophilized, redissolved in 0.1% TFA, and stored at -20°C until needed.

Results. Synthetic placental bikunin (102-159) was refolded using 20% DMSO as the oxidizing agent as described above, and purified by a 2-step purification protocol as shown below, to yield an active trypsin inhibitor (Table 1 below).

Table 1

Purification table for the isolation of synthetic placental bikunin (102-159)

TABLE 1	<del></del>		-			
Purification Step	Vol ( <b>ml)</b>	mg/ml	mg	Units <sup>C</sup> (U)	SpA (U/mg)	Yield
8.0 M Urea	4.0	3.75 a	15.0	0	0	-
20% DMSO	32.0	0.47 a	15.0	16,162	1,078	100
Kallikrein affinity	9.8	0.009 b	0.09	15,700	170,000	97
€18	3.0	0.013 ab	().04	11,964	300,000	74

<sup>a</sup>Protein determined by AAA.

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bProtein determined by OD280 nm using the extinction coefficient determined for the purified protein  $(1.7 \times 10^4 \text{ Lmol}^{-1} \text{ cm}^{-1})$ .

COne Unit is defined as the amount of material required to inhibit 50% of trypsin activity in a standard assay.

Chromatography of the crude refolded material over an immobilized bovine pancreatic kallikrein column selectively isolated 6.0% of the protein and 97% of the trypsin inhibitory activity present. Subsequent chromatography using C18 reverse-phase yielded a further purification of 2-fold, with an overall recovery of 74%. On RPHPLC, the reduced and refolded placental bikunin (102-159), exhibited elution times of 26.3 and 20.1 minutes, respectively. Mass spectroscopy analysis of the purified material revealed a molecular mass of 6829.8; a loss of 6 mass units from the starting material. This demonstrates the complete formation of the 3 disulfides predicted from the peptide sequence.

The isoelectric points of the purified, refolded synthetic placental bikunin (102-159) was determined using a Multiphor II Electrophoresis System (Pharmacia) run according to the manufacturers suggestions, together with pI standards, using a precast Ampholine PAGplate (pH 3.5 to 9.5) and focused for 1.5 hrs. After staining, the migration distance from the cathodic edge of the gel to the different protein bands was measured. The pI of each unknown was determined by using a standard curve generated by a plot of the migration distance of standards versus the corresponding pI's. With this technique, the pI of placental bikunin (102-159) was determined to be 8.3, in agreement with the value predicted from the amino acid sequence. This is lower than the value of 10.5 established for the pI of aprotinin. (Tenstad et al., 1994, Acta Physiol. Scand. 152: 33-50).

### Example 2

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### Preparation of synthetic placental bikunin (7-64)

Placental bikunin (7-64) was synthesized, refolded and purified essentially as described for placental bikunin (102-159) but with the following modifications: during refolding, the synthetic peptide was stirred for 30 hr as a solution in 20% DMSO at 25°C; purification by C18 RP-HPLC was achieved with a linear gradient of 25 to 45% acetonitrile in 0.1% TFA over 40 min (1ml/min). Active fractions from the first C18 run were reapplied to the column and fractionated with a linear gradient (60 min, 1 ml min) of 20 to 40% acetonitrile in 0.1% TFA.

25 Results. The final purified reduced peptide exhibited an MH+ = 6563, consistent with the sequence:

IHDFOLVSKV VGRORASMPF WWYNVTDGSO QLFVYGGODG MSNWYLTYEE CLYYCATV (SEQ ID MG: 4

The refolding and purification yielded a functional Kunitz domain that was active as an inhibitor of trypsin 'Table 2 below'

Table 2A

Purification table for the isolation of synthetic placental bikunin (7-64)

TABLE 2A						
Purification Step	Vol (ml)	mg/ml	mg	Units (U)	SpA (U/mg)	Yield
8.0 M Urea	8.0	2.5	20.0	0	0	•
20% DMSO	64.0	0.31	20.0	68,699	3,435	100
Kall affinity pH 4.0	11.7	0.10	1.16	43,333	36,110	62
Kall affinity pH 1.7	9.0	0.64	5.8	4972	857	72
C18-1	4.6	0.14	0.06	21,905	350,143	31.9
C18-2	1.0	0.08	0.02	7,937	466,882	11.5

The purified refolded protein exhibited an MH+ = 6558, i.e.  $5\pm1$  mass units less than for the reduced peptide. This demonstrates that refolding caused the formation of at least one appropriate disulfide bond.

The pI of placental bikunin (7-64) was determined using the methods employed to determine the pI of placental bikunin (102-159). Placental bikunin (7-64) exhibited a pI that was much higher than the predicted value (pI = 7.9). Refolded placental bikunin (7-64) migrated to the cathodic edge of the gel (pH 9.5) and an accurate pI could not be determined under these conditions.

### Continued Preparation of synthetic placental bikunin (7-64)

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Because the synthetic placental bikunin (7-64) may not have undergone complete deprotection prior to purification and refolding, refolding was repeated using protein which was certain to be completely deprotected. Placental bikunin (7-64) was synthesized, refolded and purified essentially as described for placental bikunin (102-159) but with the following modifications: during refolding, the synthetic peptide (0.27 mg/ml) was stirred for 30 hr as a solution in 20% DMSO at 25 C; purification by C18 RP-HPLC was achieved with a linear gradient of 22.5 to 50% acetonitrile in 0.1% TFA over 40 min (1 ml/min).

Results. The final purified reduced peptide exhibited an MH+=6567.5, consistent with the sequence:

IHDFCLVSKV VGRCRASMPRW WYNVTDGSC QLFVYGGCDG NSNNYLTKEE CLKKCATV (SEQ ID NO: 4)

The refolding and purification yielded a functional Kunitz domain that was as active as an inhibitor of trypsin (Table 2B below).

Table 2B
Purification table for the isolation of synthetic placental bikunin (7-64)

TABLE 2B						
Purification Step	Vol (ml)	mg/ml	mg	Units (U)	SpA (U/mg)	Yield
8.0 M Urea	4.9	2.1	10.5	0	0	
20% DMSO	39.0	0.27	10.5	236,000	22,500	100
Kallikrein Affinity (pH 2)	14.5	0.3	0.43	120,000	279,070	50.9
C18 Reverse- Phase	0.2	1.2	0.24	70,676	294,483	30.0

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The purified refolded protein exhibited an MH+ = 6561.2, i.e. 6.3 mass units less than for the reduced peptide. This demonstrates that refolding caused the formation of the expected three disulfide bonds.

The pI of refolded placental bikunin (7-64) was determined using the methods employed to determine the pI of placental bikunin (102-159). Refolded placental bikunin (7-64) exhibited a pI of 8.85, slightly higher than the predicted value (pI = 7.9).

Example 3
In vitro specificity of functional placental bikunin fragment (102-159)

Proteases. Bovine trypsin, human plasmin, and bovine pancreatic kallikrein quantitation was carried out by active site titration using prutrophenyl p'-guanidinobenzoate HCl as previously described (Chase,T., and Shaw, E., (1970) Methods Enzmol., 19: 20-27). Human kallikrein was quantitated by active site titration using bovine aprotinin as a standard and PFR-AMC as a substrate assuming a 11 complex formation. The  $K_m$  for GPK-AMC with trypsin and plasmin under the conditions used for each enzyme was 29  $\mu$ M and 726  $\mu$ M, respectively; the  $K_m$  for PFR-AMC with human plasma kallikrein and bovine pancreatic kallikrein was 457  $\mu$ M and 81.5  $\mu$ M, respectively; the  $K_m$  for AAPR-AMC with elastase was 1600  $\mu$ M. Human tissue

<sup>1948 -</sup> Charles Call Methode Edition 2012

Inhibition Kinetics: The inhibition of trypsin by placeptal bikumin 172

placental bikunin (102-159) (0-2 nM) or aprotinin (0-3 nM) in buffer A in a total volume of 1.0 ml. After 5 min. at 37°C, 15 µl of 2 mM GPK-AMC was added and the change in fluorescence (as above) was monitored. The inhibition of human plasmin by placental bikunin (102-159) and aprotinin was determined with plasmin (50 pM) and placental bikunin (102-159) (0-10 nM) or aprotinin (0-4 nM) in buffer containing 50 mM Tris-HCl (pH 7.5), 0.1 M NaCl, and 0.02% triton x-100. After 5 min. incubation at 37°C, 25 µl of 20 mM GPK-AMC was added and the change in fluorescence monitored. The inhibition of human plasma kallikrein by placental bikunin (102-159) or aprotinin was determined using kallikrein (2.5 nM) and placental bikunin (102-159) (0-3 nM) or aprotinin (0-45 nM) in 50 mM Tris-HCl (pH 8.0), 50 mM NaCl, and 0.02% triton x-100 After 5 min. at 37°C 15 µl of 20 mM PFR-AMC was added and the change in fluorescence monitored. The inhibition of bovine pancreatic kallikrein by placental bikunin (102-159) and aprotinin was determined in a similar manner with kallikrein (92 pM), placental bikunin (102-159) (0-1.6 nM) and aprotinin (0-14 pM) and a final substrate concentration of 100 μM. The apparent inhibition constant Ki\* was determined using the nonlinear regression data analysis program Enzfitter software (Biosoft, Cambridge, UK): The kinetic data from each experiment were analyzed in terms of the equation for a tight binding inhibitor:

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$$V_i/V_0 = 1 - (E_0 + I_0 + K_i^* - [(E_0 + I_0 + K_i^*)^2 - 4 E_0 I_0]^{1/2})/2E_0$$
 (2)

where  $V_i/V_0$  is the fractional enzyme activity (inhibited vs. uninhibited rate), and  $E_0$  and  $I_0$  are the total concentrations of enzyme and inhibitor, respectively. Ki values were obtained by correcting for the effect of substrate according to the equation:

$$K_i = K_i^* / (1 + [S_0] / K_m)$$
 (3)

(Boudier, C., and Bieth, J. G., (1989) Biochim Biophys Acta., 995: 36-41)

For the inhibition of human neutrophil elastase by placental bikunin (102-159) and aprotinin, elastase (19 nM) was incubated with placental bikunin (102-159) (150 nM) or aprotinin (0-7.5 µM) in buffer containing 0.1 M Tris-HCl (pH 8.0), and 0.05% triton X-100. After 5 min at 37% C, AAPM-AMC (500 µM or 1000 µM) was added and the fluorescence measured over a two-minute period Ki values were determined from Dixon plots of the form 1/V versus [I]

performed at two different substrate concentrations (Dixon et al., 1979).

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The inhibition of human tissue kallikrein by aprotinin, placental bikunin fragment (7-64) or placental bikunin fragment (102-159) was measured by the incubation of 0.35 nM human tissue kallikrein with placental bikunin (7-64) (0-40 nM) or placental bikunin (102-159) (0-2.5 nM), or aprotinin (0-0.5 nM) in a 1 ml reaction volume containing 50 mM Tris-HCl buffer pH 9.0, 50 mM NaCl, and 0.1% triton x-100. After 5 min. at 37°C, 5 ul of 2 mM PFR-AMC was added achieving 10 uM final and the change in fluorescence monitored. The Km for PFR-AMC with human tissue kallikrein under the conditions employed was 5.7 uM. The inhibition of human factor Xa (American Diagnostica, Inc, Greenwich, CT) by synthetic placental bikunin (102-159), recombinant placental bikunin, and aprotinin was measured by the incubation of 0.87 nM human factor Xa with increasing amounts of inhibitor in buffer containing 20 mM Tris (pH 7.5), 0.1 M NaCl, and 0.1% BSA. After 5 min. at 37°C, 30 ul of 20 mM LGR-AMC (Sigma) was added and the change in fluorescence monitored. The inhibition of human urokinase (Sigma) by Kunitz inhibitors was measured by the incubation of urokinase (2.7 ng) with inhibitor in a total volume of 1 ml buffer containing 50 mM Tris-HCl (pH 8.0), 50 mM NaCl, and 0.1% Triton x-100. After 5 min. at 37°C, 35 ul of 20 mM GGR-AMC (Sigma) was added and the change in fluorescence monitored. The inhibition of Factor XIa (from Enzyme Research Labs, Southbend, IN) was measured by incubating FXIa (0.1 nM) with either 0 to 800 nM placental bikunin (7-64), 0 to 140 nM placental bikunin (102-159) or 0 to 40 uM aprotinin in buffer containing 50 mM Hepes pH 7.5, 100 mM NaCl, 2 mM CaCl2, 0.01% triton x-100, and 1% BSA in a total volume of 1 ml. After 5min at 37 C, 10 ul of 40 mM Boc-Glu(OBzl)-Ala-Arg-AMC (Bachem Biosciences, King of Prussia, PA) was added and the change in fluorescence monitored.

**Results:** A direct comparison of the inhibition profiles of placental bikunin (102-159) and aprotinin was made by measuring their inhibition constants with various proteases under identical conditions. The  $K_1$  values are listed in Table 3 below.

Table 3 Ki values for the inhibition of various proteases by bikunin (102-159)

TABLE 3	<del></del>			<del></del>
Protease (concentration)	bikunin (102-159) Ki (nM)	Aprotinin Ki (nM)	Substrate (concentration)	Km (mM)
Trypsin (48.5 pM)	0.4	0.8	GPK-AMC (0.03 mM)	0.022
Chymotrypsin (5 nM)	0.24	0.86	AAPF-pNA (0.08 mM)	0.027
Bovine Pancreatic Kallikrein (92.0 pM)	0.4	0.02	PFR-AMC (0.1 mM)	0.08
Human Plasma Kallikrein (2.5 nM)	0.3	19.0	PFR-AMC (0.3 mM)	0.46
Human Plasmin (50 pM)	1.8	1.3	GPK-AMC (0.5 mM)	0.73
Human Neutrophil Elastase (19 nM)	323.0	8500.0	AAPM-AMC (1.0 μM)	1.6
Factor XIIa	>300.0	12,000.0	PFR-AMC (0.2 µM)	0.35
Human Tissue Kallikrein (0.35 nM)	0.13	0.004	PFR-AMC (10 µM)	0.0057
factor Xa (0.87 nM)	274	N.l. at 3 μM	LGR-AMC (0.6 mM)	N.D
urokinase	11000	4500	GGR-AMC (0.7 mM)	N.D.
factor XIa (0.1 nM)	15	288	E(OBz)AR-AMC (0.4 mM)	0.46

Placental bikunin (102-159) and aprotinin inhibit bovine trypsin and human plasmin to a comparable extent under the conditions employed. Aprotinin inhibited elastase with a Ki of 8.5  $\mu$ M. Placental bikunin (102-159) inhibited elastase with a Ki of 323nM. The K<sub>i</sub> value for the placental bikunin (102-159) inhibition of bovine pancreatic kallikrein was 20-fold higher than that of aprotinin inhibition. In contrast, placental bikunin (102-159) is a more potent inhibitor of human plasma kallikrein than aprotinin and binds with a 56-fold higher affinity.

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Because placental bikunin (102-159) is greater than 50 times more potent than Trasylol<sup>®</sup> as an inhibitor of kallikrein, smaller amounts of human placental bikunin, or fragments thereof (i.e. placental bikunin (102-159)) are needed than Trasylol<sup>®</sup> in order to maintain the effective patient doses of inhibitor in KIU. This reduces the cost per dose of the drug and reduces the likelihood of adverse nephrotoxic effects upon re-exposure of the medicament to patients. Furthermore, the protein is human derived, and thus much less immunogenic in man than aprotinin which is derived from cows. This results in significant reductions in the risk of incurring adverse immunologic events upon re-exposure of the medicament to patients.

### Example 4

# In vitro specificity of functional placental bikunin fragment (7-64)

In vitro specificity of functional human placental bikunin (7-64) was determined using the materials and methods as described in the Examples above.

Results: The table below shows the efficacy of placental bikurin (7-64) as an inhibitor of various serine proteases *in vitro*. Data is shown compared against data obtained for screening inhibition using either placental bikurin (102-159), or aprotinin (Trasylol<sup>®</sup>).

Table 4 A
Ki values for the inhibition of various proteases by bikunin(7-64)

Protease (concentration)	bikunin(7-64) Ki (nM)	Aprotinin Ki (nM)	bikunın (102-159 Ki (nM)	
Trypsin (48.5 pM)	0.17	0.8	0.4	
Bovine Pancreatic Kallikrein (92.0 pM)	0.4	0.02	0.4	
Human Plasma Kallikrein (2.5 nM)	2.4	19.0	0.3	
Human Plasmin (50 pM)	3.1	1.3	1.8	
Bovine chymotrypsin (5 nM)	0.6	0.9	0.2	
Factor XIIa	>300	12000	>300	
elastase	>100	8500	323	

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The results show that the amino acid sequence encoding placental bikunin (7-64) can be refolded to obtain an active serine protease inhibitor that is effective against at least four trypsin-like serine proteases.

Table 4B below also shows the efficacy of refolded placental bikunin (7-64) as an inhibitor of various serine proteases *in vitro*. Refolded placental bikunin (7-64) was prepared from protein that was certain to be completely deprotected prior to purification and refolding. Data is shown compared

Table 4B Ki values for the inhibition of various proteases by refolded bikunin (7-64)

TABLE 4B			
Protease (concentration)	bikunin (7-64) Ki (nM)	Aprotinin Ki (nM)	bikunin (102-159) Ki (nM)
Trypsin (50 pM)	0.2	0.8	0.3
Human Plasma Kallikrein (0.2 nM)	0.7	19.0	0.7
Human Plasmin (50 pM)	3.7	1.3	1.8
Factor XIIa	not done	12,000	4,500
Factor XIa (0.1 nM)	200	288	15
Human Tissue Kallikrein	2.3	0.004	0.13

Suprisingly, placental bikunin (7-64) was more potent than aprotinin at inhibiting human plasma kallikrein, and at least similar in efficacy as a plasmin inhibitor. These data show that placental bikunin (7-64) is at least as effective as aprotinin, using *in vitro* assays, and that one would expect better or similar potency *in vivo*.

Example 5

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### Expression of placental bikunin variant (102-159) in yeast

The DNA sequence encoding placental bikunin 102-159 (SEQ ID NO: 6) was generated using synthetic oligonucleotides. The final DNA product consisted (5' to 3') of 15 nucleotides from the yeast  $\alpha$ -mating factor propeptide sequence fused to the in-frame cDNA sequence encoding placental bikunin (102-159), followed by an in-frame stop codon. Upon cloning into a yeast expression vector pS604, the cDNA would direct the expression of a fusion protein comprising an N-terminal yeast  $\alpha$ -mating factor propeptide fused to the 58 amino acid sequence of placental bikunin (102-159). Processing of this fusion protein at a KEX-2 cleavage site at the junction between the  $\alpha$ -mating factor and Kunitz domain was designed to liberate the Kunitz domain at its native N-terminus.

A 5' sense oligonucleotide of the following sequence and containing a HindIII site for cloning was synthesized:

GAA GGG GTA AGC TTG GAT AAA AGA TAT GAA GAA TAC TGC ACC GCC AAC GCA GTC ACT GGG CCT TGC CGT GCA TCC TTC CCA CGC TGG TAC TTT GAC GTG GAG AGG (SEQ ID NO: 42)

A 3' antisense oligonucleotide of the following sequence and containing both a BamHI site for cloning and a stop codon was synthesized

```
530 GGA TOO OTA OTG GGG GAA GCG GAG DAT GCA GGC OTG
5 OTG AGA GCG GTA GCT GTT OTT ATT GCC CCG GCA GCC TCC ATA
GAT GAA GTT ATT GCA GGA GTT CCT CTC CAC GTC AAA GTA CCA
GCG
(SEQ ID NO: 43
```

The oligonucleotides were dissolved in 10 mM Tris buffer pH 8.0 containing 1 mM EDTA, and 12 ug of each oligo were added combined and brought to 0.25M NaCl. To hybridize, the oligonucleotides were denatured by boiling for 5 minutes and allowed to cool from 65°C to room temp over 2 hrs. Overlaps were extended using the Klenow fragment and digested with HindIII and BamHI. The resulting digested double stranded fragment was cloned into pUC19 and sequence confirmed. A clone containing the fragment of the correct sequence was digested with BamHI/HindIII to liberate the bikunin containing fragment with the following + strand sequence:

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GAA GGG GTA AGO TTG GAT AAA AGA TAT GAA GAA TAO TGO AGO GCC AAC GCA GTC ACT GGG CCT TGC CGT GCA TCC TTC CCA CGC TGG TAC TTT GAC GTG GAG AAC TCC TGC AAT AAC TTC ATC TAT GGA GGC TGC TGC GGC AAT AAG AAC AGO TAC CGC TGC GAG GAG GCC TGC ATG CGC TGC TTC CGC CAG TAG GGA TCC GEQ CGC TGC TCC CGC CAG TAG GGA TCC GEQ CGC TGC TCC CGC CAG TAG GGA TCC GEQ
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which was then gel purified and ligated into BamHI/HindIII cut pS604. The ligation mixture was extracted into phenol/chloroform and purified over a S-200 minispin column. The ligation product was directed transformed into yeast strains SC101 and WHL341 and plated on ura selection plates. Twelve colonies from each strain were re-streaked on ura drop out plates. A single colony was inoculated into 2 ml of ura DO media and grown over night at 30°C. Cells were pelleted for 2 minutes at 14000x g and the supernatants evaluated for their content of placental bikunin (102-159).

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capacity to inhibit the in titro activity of trypsin using the assay methods as described in Example 1 (1 ml assay volume). An un-used media only sample as well as a cast complete stage of pressing an inactive variant of apriting which the

negative controls. A yeast clone expressing natural aprotinin served as a positive control and is shown for comparison.

The second method to quantify placental bikunin (102-159) expression exploited use of polyclonal antibodies (pAbs) against the synthetic peptide to monitor the accumulation of the recombinant peptide using Western blots. These studies were performed only with recombinants derived from strain SC101, since these produced greater inhibitory activity than recombinants derived from strain WHL341.

To produce the pAb, two 6-8 week old New Zealand White female rabbits (Hazelton Research Labs, Denver, Pa) were immunized on day zero with 250 ug of purified reduced synthetic placental bikunin (102-159), in Complete Freund's adjuvant, followed by boosts on days 14, 35 and 56 and 77 each with 125 ug of the same antigen in Incomplete Freund's adjuvant. Antiserum used in the present studies was collected after the third boost by established procedures. Polyclonal antibodies were purified from the antiserum over protein A.

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Colonies 2.4 and 2.5 from transformation of yeast SC101 (Figure 8) as well as an aprotinin control were grown overnight in 50 ml of ura DO media at 30°C. Cells were pelleted and the supernatant concentrated 100-fold using a Centriprep 3 (Amicon, Beverly, MA) concentrator. Samples of each (30 µl) were subjected to SDS-PAGE on 10-20% tricine buffered gels (Novex, San Diego, CA) using the manufacturers procedures. Duplicate gels were either developed with a silver stain kit (Integrated Separation Systems, Nantick, MA) or transferred to nitrocellulose and developed with the purified polyclonal antibody elicited to synthetic bikunin (102-159). Alkaline-phosphatase conjugated goat anti-rabbit antibody was used as the secondary antibody according to the manufacturer's directions (Kirkegaard and Perry, Gaithersburg, MD).

# Purification of placental bikunin (102-159) from a transformed strain of SC101

Fermentation broth from a 1L culture of SC101 strain 2.4 was harvested by centrifugation (4,000 g x 30 min.) then applied to a 1.0 ml column of anhydrochymotrypsin-sepharose (Takara Biochemical Inc., CA), that was previously equilibrated with 50 mM Hepes buffer pH 7.5 containing 0.1M NaCl, 2 mM CaCl<sub>2</sub> and 0.01% (v/v) triton X-100. The column was washed with the same buffer but containing 1.0 M NaCl until the A280nm declined to zero, whereupon the column was eluted with 0.1M formic acid pH 2.5. Eluted fractions were pooled and applied to a C18 column (Vydac, 5um, 4.6 x 250 mm)

previously equilibrated with 0.1% TFA, and eluted with a 50 min. linear gradient of 20 to 80% acetonitrile in 0.1% TFA. Fractions containing placental bikunin (102-159) were pooled and re-chromatographed on C18 employing elution with a linear 22.5 to 50% acetonitrile gradient in 0.1% TFA.

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Results. Figure 8 shows the percent trypsin activity inhibited by twelve colonies derived from the transformation of each of strains SC101 and WHL341. The results show that all twelve colonies of yeast strain SC101 transformed with the trypsin inhibitor placental bikunin (102-159) had the ability to produce a substantial amount of trypsin inhibitory activity compared to the negative controls both of which showed no ability to inhibit trypsin. The activity is therefore related to the expression of a specific inhibitor in the placental bikunin variant (102-159) transformed cells. The yeast WHL341 samples contained minimal trypsin inhibitory activity. This may be correlated to the slow growth observed with this strain under the conditions employed.

Figure 9 shows the SDS-PAGE and western analysis of the yeast SC101 supernatants. Silver stained SDS-PAGE of supernatants derived from recombinant yeasts 2.4 and 2.5 expressing placental bikunin (102-159) as well as from the yeast expressing aprotinin yielded a protein band running at approximated 6 kDa, corresponding to the size expected for each recombinant Kunitz inhibitor domain. Western analysis showed that the 6 kDa bands expressed by stains 2.4. and 2.5 reacted with the pAb elicited to placental bikunin (102-159). The same 6 kDa band in the aprotinin control did not react with the same antibody, demonstrating the specificity of the antibody for the placental bikunin variant (102-159).

The final preparation of placental bikunin C-terminal domain was highly pure by silver-stained SDS-PAGE (Figure 10). The overall recovery of broth-derived trypsin inhibitory activity in the final preparation was 31%. N-terminal sequencing of the purified inhibitor indicated that 40% of the protein is correctly processed to yield the correct N-terminus for placental bikunin (102-159) while about 60 % of the material contained a portion of the yeast α-mating

ranirien.

In conclusion, the accumulation both of a protease inhibitor activity and a protein immunochemically related to synthetic bikings 102-159 in termination of the conclusion which is the solution of placenta now account of the concepts.

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one of the transformed lines provided proof of expression of placental bikunin in the recombinant yeast strains described herein, showing for the first time the utility of yeasts for the production of placental bikunin fragments.

Additional constructs were prepared in an effort to augment the expression level of the Kunitz domain contained within placental bikunin 102-159, as well as to increase the yield of protein with the correct N-terminus. We hypothesized that the N-terminal residues of placental bikunin 102-159 (YEEY--) may have presented a cleavage site that is only poorly recognized by the yeast KEX-2 protease that enzymically removes the yeast a-factor pro-region. Therefore, we prepared yeast expression constructs for the production of 10 placental bikunin 103-159 (N-terminus of EEY...), 101-159 (N-terminus of NYEEY...) and 98-159 (DMFNYEEY...) in order to modify the P' subsites surrounding the KEX-2 cleavage site. To attempt to augment the levels of recombinant protein expression, we also used the yeast preferred codons rather than mammalian preferred codons in preparing some of the constructs 15 described below. The constructs were essentially prepared as described above for placental bikunin 102-159 (defined as construct #1) but with the following modifications:

20 Construct #2 placental bikunin 103-159, yeast codon usage A 5' sense oligonucleotide

and 3' antisense oligonucleotide

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ACTGGATCCT CATTGGCGAA AACATCTCAA CATACAGGCT

TCTTCAGATC TGTAAGAATT TTTATTACCT CTACAACCAC
CGTAAATAAA ATTATTACAA GAATTTCTTT CAACATCAAA
GTACCATCT (SEQ ID NO: 56)

were manipulated as described for the production of an expression construct (construct #1 above) for the expression of placental bikunin 102-159

Construct #3 placental bikurun 101-159, yeast codon usage

A 5' sense oligonucleotide

GAAGGGGTAA GCTTGGATAA AAGAAATTAC GAAGAATACT GTACTGCTAA TGCTGTTACT GGTCCATGTA GAGCTTCTTT TCCAAGATGGTACTTTGATGTTGAAAGA (SEQ ID NO: 57)

and the same 3' antisense oligonucleotide as used for construct #2, were manipulated as described for the production of an expression construct (construct #1 above) for the expression of placental bikunin 102-159.

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Construct #4 placental bikunin 98-159, yeast codon usage A 5' sense oligonucleotide

GAAGGGGTAA GCTTGGATAA AAGAGATATG TTTAATTACG 15 AAGAATACTG TACTGCTAAT GCTGTTACTG GTCCATGTAG AGCTTCTTTT CCAAGATGGT ACTTTGATGT TGAAAGA (SEQ ID NO: 58)

and the same 3' antisense oligonucleotide as used for construct #2. were manipulated as described for the production of an expression construct (construct #1 above).

Yeast strain SC101 (MATa, ura 3-52, suc 2) was transformed with the plasmids containing each of the above cDNAs, and proteins were expressed using the methods that were described above for the production of placental bikunin 102-159 with human codon usage. Approximately 250 ml of each yeast culture was harvested, and the supernatant from centrifugation (15 min x 3000 RPM) separately subjected to purification over 1 ml columns of kailikrein-sepharose as described above. The relative amount of trypsin inhibitory activity in the applysate, the amount of purified protein recovered and the N-terminal sequence of the purified protein were determined and are listed below in Table

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Table 7
Relative production levels of different proteins containing the C-terminal Kunitz domain of placental bikunin

TABL	LE 7				
Cons	truct	Relative conc. of inhibitor in applysate	N-terminal amount (pmol)	sequencing: sequence	Comments
#2	103-159	none detected	none	none	no expression
#3	101-159	25 % inhibition	none	none	low expression
	98-159 ession ect product	93 % inhibition	910	DMFNYE-	good
#1	102-159	82 % inhibition	480	AKEEGV-	expression of active incorrectly processed protein

The results show that placental bikunin fragments of different lengths that contain the C-terminal Kunitz domain show wide variation in capacity to express functional secreted protein. Constructs expressing fragments 101-159 and 103-159 yielded little or low enzymic activity in the supernatants prior to purification, and N-terminal sequencing of 0.05 ml aliquots of each purified fraction yielded undetectable amounts of inhibitor. On the other hand expression either of placental bikunin 102-159 or 98-159 yielded significant amounts of protease activity prior to purification. N-terminal sequencing however showed that the purified protein recovered from expression of 102-159 was once again largely incorrectly processed, exhibiting an N-terminus consistent with processing of the majority of the pre-protein at a site within the yeast α-mating factor pro-sequence. The purified protein recovered from expression of placental bikunin 98-159 however was processed entirely at the correct site to yield the correct N-terminus. Furthermore, nearly twice as much protein was recovered as compared to the recovery of placental bikumin 102-159. Placental bikunin 98-159 thus represents a preferred fragment length for the production of the C-terminal Kunitz domain of placental bikunin by the  $\alpha$ mating factor pre-pro sequence/ KEX-2 processing system of S. cerevisiae,

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### Example 6

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### Alternative procedure for yeast expression

The 58 amino acid peptide derived from the R74593 translation product can also be PCR amplified from either the R87894-R74593 PCR product cloned into the TA vector<sup>TM</sup> (Invitrogen, San Diego, CA) after DNA sequencing or from human placental cDNA. The amplified DNA product will consist of 19 nucleotides from the yeast  $\alpha$ -matting factor leader sequence mated to the R74593 sequence which codes for the YEEY--CFRQ (58 residues) so as to make the translation product in frame, constructing an  $\alpha$  -mating factor/Kunitz domain fusion protein. The protein sequence also contains a kex 2 cleavage which will liberate the Kunitz domain at its native N-terminus.

The 5' sense oligonucleotide which contains a HindIII site for cloning will contain the following sequence:

GCCAAGCTTG GATAAAAGAT ATGAAGAAT ACTGCACCGC CAACGCA 15 (SEQ ID NO: 30

The 3' antisense oligonucleotide contains a BamHI site for cloning as well as a stop codon and is of the following sequence:

GGGGATCCTC ACTGCTGGCG GAAGCAGCGG AGCAT (SEQ ID NO: 31 20

The full 206 nucleotide cDNA sequence to be cloned into the yeast expression vector is of the following sequence:

DOAAGOTTGG ATAAAAGATA TGAAGAATAO TGOADDGCGA ACGODGAA 25 TGGGCCTTGC CGTGCATCCT TCCCACGCTG GTACTTTGAC GTCGACAGGA ACTOOTGOAA TAACTTOATO TATGGAGGOT GCCGGGGGCAA TAAGAAGAGG TACOGOTOTO AGGAGGOOTO CATGOTOCGO TGOTTOCGOO AGGAGTGAGO ATCCCC (SEQ ID NO: 32)

After PCR amplification, this DNA will be digested with HindIII. BamHI and clamped into the upact princes in . . -the mattre

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using the methods described in US patent 5,164,482. The URA 3- yeast 35 transformants are isolated and cultivated under inducing conditions. The yield t res nebinant Placental bikunun sarlantsus determine puolitiks juoliteis

amount of trypsin inhibitory activity that accumulated in the culture supernatants over time using the in vitro assay method described above. Fermentation broths are centrifuged at 9000 rpm for 30 minutes. supernatant is then filtered through a 0.4 then a 0.2 µm filter, diluted to a conductivity of 7.5 ms, and adjusted to pH 3 with citric acid. The sample is then batch absorbed onto 200 ml of S-sepharose fast flow (Pharmacia) in 50 mM sodium citrate pH 3 and stirred for 60 min. The gel is subsequently washed sequentially with 2 L of each of: 50 mM sodium citrate pH 3.0; 50 mM Tris-HCL pH 9.0; 20 mM HEPES pH 6.0. The washed gel is transferred into a suitable column and eluted with a linear gradient of 0 to 1 M sodium chloride in 20 mM HEPES pH 6.0. Eluted fractions containing in vitro trypsin inhibitory activity are then pooled and further purified either by a) chromatography over a column of immobilized anhydrotrypsin (essentially as described in Example 2); b) by chromatography over a column of immobilized bovine kallikrein; or c) a combination of conventional chromatographic steps including gel filtration and/or anion-exchange chromatography.

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# Example 7 Isolation and characterization of native human placental bikunin from placenta

Bikunin protein was purified to apparent homogeniety from whole frozen placenta (Analytical Biological Services, Inc, Wilmington, DE). The placenta (740 gm) was thawed to room temperature and cut into 0.5 to 1.0 cm pieces, placed on ice and washed with 600 ml PBS buffer. The wash was decanted and 240 ml of placenta pieces placed into a Waring blender. After adding 300 ml of buffer consisting of 0.1 M Tris (pH 8.0), and 0.1 M NaCl , the mixture was blended on high speed for 2 min, decanted into 750.0 ml centrifuge tubes, and placed on ice. This procedure was repeated until all material was processed. The combined slurry was centrifuged at 4500 x g for 60 minutes at 4°C. The supernatant was filtered through cheese cloth and the placental bikunin purified using a kallikrein affinity column made by covalently attaching 70 mg of bovine pancreatic kallikrein (Bayer AG) to 5.0 mls of CNBr activated Sepharose (Pharmacia) according to manufacturers instruction. The material was loaded onto the affinity column at a flow rate of 2.0 ml/min and washed with 0.1 M Tris (pH 8.0), 0.1 M NaCl until absorbance at 280 nm of the wash could no longer be detected. The column was further washed with 0.1 M Tris (pH 8.0), 0.5 M NaCl and then eluted with 3 volumes of 0.2 M acetic acid,

pH 4.0. Fractions containing kallikrein and trypsin inhibitory (see below) activity were pooled, frozen, and lyophilized. Placental bikunin was further purified by gel-filtration chromatography using a Superdex 75 10/30 (Pharmacia) column attached to a Beckman System Gold HPLC system. Briefly, the column was equilibrated in 0.1 M Tris, 0.15 M NaCl, and 0.1% Triton X-100 at a flow rate of 0.5 ml/min. The lyophilized sample was reconstituted in 1.0 ml of 0.1 M Tris, pH 8.0 and injected onto the gel-filtration column in 200 μl aliquots. Fractions were collected (0.5 ml) and assayed for trypsin and kallikrein inhibitory activity. Active fractions were pooled, and the pH of the solution adjusted to 2.5 by addition of TFA. The material was directly applied to a Vydac C18 reverse-phase column (5 micron, 0.46 x 25 cm) which had been equilibrated in 20% acetonitrile in 0.1 %TFA. Separation was achieved using a linear gradient of 20 to 80% acetonitrile in 0.1% TFA at 1.0 ml/min over 50minutes after an initial 20 minute wash at 20% acetonitrile in 0.1% TFA. Fractions (1ml) were collected and assayed for trypsin and kallikrein inhibitory activity. Fractions containing inhibitory activity were concentrated using a speed-vac concentrator (Savant) and subjected to N-terminal sequence analysis.

## Functional assays for Placental Bikunin:

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Identification of functional placental bikunin was achieved by measuring its ability to inhibit bovine trypsin and human plasma kallikrein. Trypsin inhibitory activity was performed in assay buffer (50 mM Hepes, pH 7.5, 0.1 M NaCl, 2.0 mM CaCl2, 0.1% Triton x-100) at room temperature in a 96-well microtiter plate (Perkin Elmer) using Gly-Pro-Lys-Aminomethylcoumarin as a substrate. The amount of coumarin produced by trypsin was determined by measuring the fluorescence (ex = 370 nm, em = 432 nm) on a Perkin-Elmer LS-50B fluorimeter equipped with a plate reader. Trypsin (23 μg in 100 μl buffer) was mixed with 20 μl of the sample to be tested and incubated for 10 minutes at 25°C. The reaction was started by the addition of 50 μl of the substrate GPK-AMC (33 μM final) in assay buffer. The fluorescence intensity was measured and the % inhibition for each fraction was determined by

where Fo is the fluorescence of the unknown and FI is the fluorescence of the trypsin only control. Kallikrein inhibitory activity of the fractions was similarly teasured using 50 pM Falderson assay buffer. 50 pM. Tris. ( H. 50 p. M.

NaCl,  $0.1^{\rm o}_{\rm o}$  triton x-100) and  $66.0\,\mu M$  Pro-Phe-Arg-AMC as a substrate.

# Determination of the in vitro specificity of placental bikunin

The *In vitro* specificity of native human placental bikunin was determined using the materials and methods as described in the preceding examples above. Placental bikunin was quantified by active site titration against a known concentration of trypsin using GPK-AMC as a substrate to monitor the fraction of unbound trypsin.

### 10 Protein Sequencing

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The 1 ml fraction (C18-29 Delaria) was reduced to 300 ml in volume, on a Speed Vac, to reduce the amount of organic solvent. The sample was then loaded onto a Hewlett-Packard miniature biphasic reaction column, and washed with 1 ml of 2% trifluoroacetic acid. The sample was sequenced on a Hewlett-Packard Model G1005A protein sequencing system using Edman degradation. Version 3.0 sequencing methods and all reagents were supplied by Hewlett-Packard. Sequence was confirmed for 50 cycles.

Results. Placental Bikunin was purified to apparent homogeniety by sequential kallikrein affinity, gel-filtration, and reverse-phase chromatography (see purification table below):

Table 5 Purification table for native Placental Bikunin (1-179)

TABLE 5							
Step	Vol (ml)	OD (/ml)	280	OD 280	Units <sup>a</sup> (U)	Units/OD 280	
Placenta Supernatant	1800.0	41.7		75,060	3,000,000	40.0	
Kallikrein Affinity pH 4.0	20.0	0.17		3.36	16,000	4,880	
Kallikrein Affinity pH 1.7	10.2	0.45		4.56	12.000	2,630	
Superdex 75	15.0	0.0085		0.13	3.191	24,546	

<sup>&</sup>lt;sup>a</sup>One Unit is defined as that amount which inhibits 50% of trypsin activity in a standard assay.

The majority of the kallikrein and trypsin inhibitory activity eluted from the kallikrein affinity column in the pH 4.0 elution. Subsequent gel-filtration chromatography (Figure 5) yielded a peak of kallikrein and trypsin inhibitory

activity with a molecular weight range of 10 to 40 kDa as judged by a standard curve generated by running molecular weight standards under identical conditions. Reverse-phase C18 chromatography (Figure 6) yielded 4 peaks of inhibitory activity with the most potent—eluting at approximately 30 % acetonitrile. The activity associated with the first peak to elute from C18 (fraction 29) exhibited an amino acid sequence starting with amino acid 1 of the predicted amino acid sequence of placental bikunin (ADRER...; SEQ ID NO: 1), and was identical to the predicted sequence for 50 cycles of sequencing (underlined amino acids in Figure 3). Cysteine residues within this sequence stretch were silent as expected for sequencing of oxidized protein. The cysteine residues at amino acid positions 11 and 20 of mature placental bikunin were later identified from sequencing of the S-pyridylethylated protein whereupon PTH-pyridylethyl-cysteine was recovered at cycles 11 and 20.

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Interestingly, the asparagine at amino acid residue number 30 of the sequence (Figure 3) was silent showing that this site is likely to be glycosylated. Fraction 29 yielded one major sequence corresponding to that of placental bikunin starting at residue #1 (27 pmol at cycle 1) plus a minor sequence (2 pmol) also derived from placental bikunin starting at residue 6 (SIHD...). This shows that the final preparation sequenced in fraction 29 is highly pure, and most likely responsible for the protease inhibitory activity associated with this fraction (Figure 6).

Accordingly, the final preparation of placental bikunin from C18 chromatography was highly pure based on a silver-stained SDS-PAGE analysis (Figure 7), where the protein migrated with an apparent Mr of 24 kDa on a 10 to 20 % acrylamide tricine gel (Novex, San Diego, CA) calibrated with the following molecular weight markers: insulin (2.9 kDa); bovine trypsin inhibitor (5.8 kDa); lysozyme (14.7 kDa);  $\beta$ -lactaglobulin (18.4 kDa), carbonic anhydrase (29 kDa); and ovalbumin (43 kDa). The above size of placental bikunin on SDS-PAGE is consistent with that predicted from the full length coding sequence (Figure 4F).

As expected based on the N-terminal sequencing results described

the purified preparation detected on gels by silver stain (Figure 7). However when the same preparation was reacted with an antibody elicited to synthetic placental bik unity (102-159), a band corresponding to the full length protect was a pseconal feature a tragment may a magrated with sea that the contraction of t

159) of approximately 6 kDa was observed. The simplest interpretation of these results is that the purified preparation had undergone degradation subsequent to purification to yield an N-terminal fragment comprising the N-terminal domain and a C-terminal fragment comprising the C-terminal domain. Assuming that the fragment reactive against antiserum to placental bikunin (7-64) is devoid of the C-terminal end of the full length protein, the size (24 kDa) would suggest a high state of glycosylation.

Table 6. below shows the potency of *in vitro* inhibition of various serine proteases by placental bikunin. Data are compared with that obtained with aprotinin (Trasylol $^{\textcircled{\$}}$ ).

Table 6
Ki values for the inhibition of various proteases by placental bikunin

TABLE 6			
Protease (concentration)	Placental Bikunin Ki (nM)	Aprotinin Ki (nM)	
Trypsin (48.5 pM)	0.13	0.8	
Human Plasmin (50 pM)	1.9	13	

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The results show that placental bikunin isolated from a natural source (human placenta) is a potent inhibitor of trypsin-like serine proteases.

### Example 8

# 20 Expression pattern of placental bikunin amongst different human organs and tissues

A multiple tissue northern was purchased from Clontech which contained 2 µg of polyA+ RNA from human heart, brain, placenta, lung, liver, skeletal muscle, kidney, and pancreas. Two different cDNA probes were used: 1) a gel purified cDNA encoding placental bikunin (102-159); 2) the 780 base pair PCR-derived cDNA (Figure 4E) liberated from a TA clone by digestion with EcoRI and gel purified. Each probe was labeled using <sup>32</sup>P-dCTP and a random priming labeling kit from Boehringer Mannheim Biochemicals (Indiana), then used to hybridize to the multiple tissue northern according to the manufacturers specifications. Autoradiographs were generated using Biomax film with an 18 hr exposure time, and developed using a Umax Scanner and scanned using Adobe Photoshop.

Results. The pattern of tissue expression observed using a placental bikumin (102-159) probe (Figure 11A) or a larger probe containing both Kumitz domains of placental bikumin (Figure 11B) was essentially the same as might be expected. The placental bikumin mRNA was most abundant in pancreas and placenta. Significant levels were also observed in lung, brain and kidney, while lower levels were observed in heart and liver, and the mRNA was undetectable in skeletal muscle. The transcript size was 1.95 kilobases in all cases, in close agreement with the predicted size of placental bikumin deduced both from EST overlay and cloning of full length cDNA described in preceding sections.

The broad tissue distribution of the mRNA shows that placental bikunin is broadly expressed. Since the protein also contains a leader sequence it would have ample exposure to the human immune system, requiring that it become recognized as a self protein. Additional evidence for a broad tissue distribution of placental bikunin mRNA expression was derived from the fact that some of the EST entries with homology to placental bikunin (Figure 4B) were derived from human adult and infant brain, and human retina, breast, ovary, olfactory epithelium, and placenta. It is concluded therefore that administration of the native human protein to human patients would be unlikely to elicit an immune response.

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Interestingly, the expression pattern of placental bikunin is somewhat reminiscent of that for bovine aprotinin which is found in high levels in bovine lung and pancreas. To further elucidate the expression pattern of placental bikunin, RT-PCR of total RNA from the following human cells was determined: un-stimulated human umbilical vein endothelial cells (HUVECs), HK-2 (line derived from kidney proximal tubule), TF-1 (erythroleukemia line) and phorbolester (PMA)-stimulated human peripheral blood leukocytes. The probes used:

CACCTGATCGCGAGACCCC (sense; SEQ ID NO: 59); CTGGCGGAAGCAGCGGAGCATGC (antisense; SEQ ID NO: 60),

and any an 800 p.p. actin tragment. Whereas the 800 p p tragment identified on agarose gels with ethidium bromide was of equal intensity in all lanes, the 600 b p. placental bikunin frazment was absent from the HUVEC above processor against an early and the first time their enumes. We are underparted as the first continues.

bikumin is not expressed in at least some endothelial cells but is expressed in some leukocyte populations.

### Example 9

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Purification and properties of Placental Bikunin (1-170) highly purified from a Baculovirus / Sf9 expression system

A large fragment of Placental bikunin containing both Kunitz domains (Placental Bikunin 1-170) was expressed in Sf9 cells as follows. Placental bikunin cDNA obtained by PCR (Figure 4E) and contained within a TA vector (see previous Examples) was liberated by digestion with HindIII and Xba1 yielding a fragment flanked by a 5' XbaI site and 3' HindIII site. This fragment was gel purified and then cloned into the M13mp19 vector (New England Biolabs, Beverly, MA). In vitro mutagenesis (Kunkel T.A., (1985) Proc. Natl. Acad. Sci. USA, 82: 488-492) was used to generate a Pst1 site 3' to the XbaI site at the 5' end, but 5' to the sequence encoding the ATG start site, natural placental bikunin signal peptide and mature placental bikunin coding sequence. The oligonucleotide used for the mutagenesis had the sequence:

5' CGC GTC TCG GCT GAC CTG GCC CTG CAG ATG GCG CAC GTG TGC 20 GGG 3' (SEQ ID NO: 61)

A stop codon (TAG) and BgIII / XmaI site was similarly engineered at the 3' end of the cDNA using the oligonucleotide:

25 5' CTG CCC CTT GGC TCA AAG TAG GAA GAT CTT CCC CCC GGG GGG GTG GTT CTG GCG GGG CTG 3' (SEQ ID NO: 62).

The stop codon was in frame with the sequence encoding placental bikunin and caused termination immediately following the Lysine at amino acid residue 170, thus encoding a truncated placental bikunin fragment devoid of the putative transmembrane domain. The product from digestion with Pstl and BglII was isolated and cloned into the BacPac8 vector for expression of Placental bikunin fragment (1-170) which contains both Kunitz domains but which is truncated immediately N-terminal to the putative transmembrane segment.

The expression of Bikunin by Sf-9 insect cells was optimal at a multiplicity of infection of 1 to 1 when the medium was harvested at 72 h post

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infection. After harvesting, the baculovirus cell culture supernatant (2L) was adjusted to pH 8.0 by the addition of Tris-HCl. Bikunin was purified by chromatography using a 5 ml bovine pancreatic kallikrein affinity column as previously described in Example 7 for the purification of native placental bikunin from placenta. Eluted material was adjusted to pH 2.5 with TFA and subjected to chromatography on a C18 reverse-phase column (1.0 x 25 cm) equilibrated in 10% acetonitrile in 0.1% TFA at a flow rate of 1 ml/min. The bikunin was eluted with a linear gradient of 10 to 80% acetonitrile in 0.1% TFA over 40 min. Active fractions were pooled, lyophilized, redissolved in 50 mM Hepes (pH 7.5), 0.1 M NaCl, 2 mM CaCl2, and 0.1% triton x-100, and stored at -20°C until needed. The concentration of recombinant bikunin was determined by amino acid analysis.

Results. Recombinant bikunin was purified from baculovirus cell culture supernatant using a 2-step purification protocol as shown below, to yield an active trypsin inhibitor (Table 8 below).

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Table 8
Purification of recombinant bikunin from transformed culture supernatant

TABLE 8					
Purification Step	Vol (ml)	OD 280/ml	OD 280 total	Units (U)	Specific activity (U/OD)
Supernatant	2300.0	9.0	20,7(0)	6.150,000	297
Kallikrein affinity	23.0	0.12	2.76	40,700	14.746
C18 reverse-phase	0.4	3.84	1.54	11.111	72,150

Chromatography of the crude material over an immobilized bovine pancreatic kallikrein affinity column selectively isolated 0.013 % of the protein and 0.67 % of the trypsin inhibitory activity present. The majority of the trypsin inhibitory activity present in the starting supernatant did not pind to the immobilized kallikrein and is not related to bikunin cresults not shown

immunoblots to rabbit anti-placental bikurun 102-159 (not shown). N-terminal sequencing (26 typles) yielded the expected sequence for margin placental section. The property of the expected sequence for margin placents.

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peptide was correctly processed in Sf9 cells.

Purified placental bikunin from Sf9 cells (100 pmol) was pyridylethylalkylated, CNBr digested and then sequenced without resolution of the resulting fragments. Sequencing for 20 cycles yielded the following N-terminii:

	Sequence	Amount	Placental bikunin residue #
10	LRCFrQQENPP-PLG ADRERSIHDFCLVSKVVGRC FNYeEYCTANAVTGPCRASF PrY-V-dGS-Q-F-Y-G	21 pmol 20 pmol 16 pmol 6 pmol	154 - 168 (SEQ ID NO: 63) 1 - 20 (SEQ ID NO: 64) 100 - 119 (SEQ ID NO: 65) 25 - 43 (SEQ ID NO: 66)

Thus N-terminii corresponding to each of the expected four fragments were recovered. This confirms that the Sf9 expressed protein contained the entire ectodomain sequence of placental bikunin (1-170). N-terminal sequencing (50 cycles) of an additional sample of undigested Placental Bikunin (1-170) resulted in an amino acid sequence which at cycle 30 was devoid of any PTH-amino acid (PTH-asparagine was expected). A similar result was obtained upon sequencing of the natural protein from human placenta (Example 7) and is consistent with this residue being glycosylated as predicted from the amino acid sequence surrounding this asparagine residue. Furthermore, the cysteine residues within this region were also silent consistent with their participation in disulfide bonding.

# 25 Example 10

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# Inhibition specificity of purified placental bikunin derived from Sf9 cells.

The *in vitro* specificity of recombinant bikunin was determined using the materials and methods as described in Examples 3, 4 and 7. In addition, the inhibition of human tissue kallikrein by bikunin was measured by the incubation of 0.35 nM human tissue kallikrein recombinant bikunin in buffer containing 50 mM Tris (pH 9.0), 50 mM NaCl, and 0.01% triton x-100. After 5 min. at  $37^{\circ}$ C, 5  $\mu$ l of 2 mM PFR-AMC was added and the change in fluorescence monitored.

Inhibition of tissue plasminogen activator (tPA) was also determined as follows: tPA (single chain form from human melanoma cell culture from Sigma Chemical Co, St Louis, MO) was pre-incubated with inhibitor for 2 hr at room temperature in 20 mM Tris buffer pH 7.2 containing 150 mM NaCl, and 0.02% sodium azide. Reactions were subsequently initiated by transfer to a reaction

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system comprising the following initial component concentrations: tPA (7.5 nM), inhibitor 0 to 6.6  $\mu$ M, DIle-Lpro-Larg-pNitroaniline (1mM) in 28 mM Tris buffer pH 8.5 containing 0.004 % (v/v) triton x-100 and 0.005% (v/v) sodium azide. Formation of p-Nitroaniline was determined from the A405nm measured following incubation at 37 C for 2hr.

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The table below show the efficacy of recombinant bikunin as an inhibitor of various serine proteases *in vitro*. Data is shown compared against data obtained for screening inhibition using either recombinant bikunin, or aprotinin.

Table 9
Comparisons of Ki values for the inhibition of various proteases by recombinant placental bikunin (1-170) or aprotinin

TABLE 9		
Protease (concentration)	Recombinant Bikunin Ki (nM)	Aprotinin Ki (nM)
Trypsin (48.5 pM)	0.064	0.8
Human Plasma Kallikrein (2.5 nM)	0.18	19.0
Human Tissue Kallikrein (0.35 nM)	0.04	0.004
Bovine Pancreatic Kallikrein (100 pM)	0.12	0.02
Human Plasmin (50 pM)	0.23	1.3
factor Xa (0.87 nM)	180	5% Inhibition at 31 IJM
factor XIa (0.1 nM)	3.0	288
tissue plasminogen activator (7.5 nM)	< 60)	no inhibition at 6.6 $\mu$ M
Tissue Factor VIIa	800	no inhibition at 1 UM

The results show that recombinant bikunin can be expressed in insect cells to yield an active protease inhibitor that is effective against at least five different serine protease inhibitors. Recombinant bikunin was more potent than aprotinin against human plasma kallikrein, trypsin and plasmin. Surprisingly, the recombinant bikunin was more potent that the synthetically derived below a fragment. This are followed to the potent and the synthetically derived.

Besides measuring the potencies against specific proteases, the capacity of placental bikunin (1-170) to prolong the activated partial thromboplastin

and the second second second second second

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aprotinin. Inhibitor was diluted in 20 mM Tris buffer pH 7.2 containing 150 mM NaCl and 0.02% sodium azide and added (0.1 ml) to a cuvette contained within an MLA Electra 800 Automatic Coagulation Timer coagulometer (Medical Laboratory Automation, Inc., Pleasantville, N.Y.). The instrument was set to APTT mode with a 300 sec. activation time and the duplicate mode. Following addition of 0.1 ml of plasma (Specialty Assayed Reference Plasma lot 1-6-5185, Helena Laboratories, Beaumont, TX), the APTT reagent (Automated APTT-lot 102345, from Organon Teknika Corp., Durhan, NC) and 25 mM CaCl2 were automatically dispensed to initiate clotting, and the clotting time was monitored automatically. The results (Figure 14) showed that a doubling of the clotting time required approximately 2  $\mu$ M final aprotinin, but only 0.3  $\mu$ M Sf9 derived placental bikunin. These data show that placental bikunin is an effective anticoagulant, and usefull as a medicament for diseases involving pathologic activation of the intrinsic pathway of coagulation.

Although certain embodiments of the invention have been described in detail for the purpose of illustration, it will be readily apparent to those skilled in the art that the methods and formulations described herein may be modified without departing from the spirit and scope of the invention. Accordingly, the invention is not limited except as by the appended claims.

#### WE CLAIM:

1. A substantially purified protein, having serine protease inhibitory activity, selected from the group of proteins consisting of materials each of which comprises one of the following amino acid sequences, the amino acids of said sequences being numbered in accordance with the amino acid sequence of native human placental bikunin shown in figure 4F in which the N-terminal residue generated by removal of signal peptide is designated as residue 1:

```
10
              ADRERSIHDF CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN
                                                                                                                                                                                                       50
              YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRO DSEDHSSDMF
                                                                                                                                                                                                     100
              NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE
                                                                                                                                                                                                    150
               ACMLRCFRQQ ENPPLPLGSK
                                                                                                                                                                                                     170
               (SEQ ID NO: 52);
15
              MAGLOGI RRSRAFLALL GSLILSGVLA
                                                                                                                                                                                                       -:
              ADRERSIHDF CLVSKVVGRC RASMPRWWYN VTDGSCQLFV YGGCDGNSNN
                                                                                                                                                                                                       50
              YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF
                                                                                                                                                                                                    100
              NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE
                                                                                                                                                                                                     150
20 ACMLRCFRQQ ENPPLPLGSK VVVLAGLFVM VLILFLGASM VYLIRVARRN
                                                                                                                                                                                                     200
               QERALRIVWS SGDDKEOLVK NTYVL
                                                                                                                                                                                                     225
               (SEQ ID NO: 49);
               ADRERSIHDE CLUSKVUGRC RASMPRWWYN UTDGSCQLEU YGGCDGNSNN
                                                                                                                                                                                                      5.3
25
             YLTYEECLKK CATUTENATG CLATSRNAAD SSVPSAPPRO DSEDHSSDMF
                                                                                                                                                                                                     :0:
               MYEEYCTANA UTGPORASEP RWYFDVERNS CHNFIYGGOR GNYMSYRSEE
                                                                                                                                                                                                     150
               ACMLROFRQQ ENPPLPLGSK WWWLAGLFVM WLILFLGASM WYLIRWARRN
                                                                                                                                                                                                     200
               QERALRTVWS SGDDKEQLVK NTYVL
                                                                                                                                                                                                     225
               SEQ ID No: Tol;
 30
                          AGSFLAWL GSLLLSGVLA -1
                sentenatione organizatione dangerens campararem . Ancesagen
                ores. CAMP of the Application of the contraction of
 35
               ACMIRCFRQQ ENPPLPIGSK VVVLAGAVS
                                                                                                                                                                                                     : 7 3
                  SEQ ID NO: 2 :
```

	MLR AEADGVERLU GELLUSGVLA	٠.
	ADRERSIHDE CLYSKYTGRO RASMPRWWYN YTDGSCQLEV YGGCDGNSNN	5 :
	YLTKEECLKK CATVTENATG DLATSRNAAD SSVESAPERQ DSECHSSOME	111
	NYEEYOTANA VTGPORASEP RWYEDVERNS CNNFIYGGOR GNKNSVRSEE	150
5	ACMERCFPQQ ENPPLPEGSK VVVEAGLFVM VEILFEGASM VYLIFVARRN	200
	QERALRIVWS SGDDKEQUVK NIYVL	225
	(SEQ ID NO: 45);	
	MAQLCGL RRSRAFLALL GSLLLSGVLA	- 1
10	ADRERSIHDE CLYSKWYGRO RASMPEWWYN VTDGSCOLEV YGGCDGNSUN	5 0
	YLTKEECLKK CATVTENATG DLATSPNAAD SSVPSAPRPQ DSEDHSSDMF	100
	NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE	150
	ACMLROFRQQ ENPPLPLGSK VVVLAGLFVM VLILFLGASM VYLIFVAPPN	200
	QERALRTUWS FGD	213
15	(SEQ ID NO: 47):	
	ADRERSIHDE CLYSKY/GRO RASMPEWWYN VTDGSCQLEV YGGODGNStNI	5.0
	YLTKEECLKK CATVTENATG DLATSRNAAD SSVPSAPPRQ DSEDHSSDMF	100
	NYEEYCTANA VTGPCRASFP RWYFDVERNS CNNFIYGGCR GNKNSYRSEE	150
20	ACMLRCFRQQ ENPPLPLGSK VVVLAGLFVM VLILFLGASM VYLIEVARRN	200
	QERALRTVWS FGD	213
	(SEQ ID NO: 71);	
	IHDF CLVSKVVGRC EASMPRWWYN VTDGSCQLFV VGGCCGNSNN 50	
25	YLTKEECLKK CATV 64	
	(SEQ ID No: 4);	
	CLVSKVVGRC PASMPRWWYN VTDGSCQLFV YGGCDGNSNN 50	
	YLTKEECLEK C 61	
30	(SEQ ID No: 5);	
	YEEYOTANA VIGPORASER PWYFDVERNS CHNFIYGGOF GMYNSYFSEE	15:
	ACMLRCFRQ	159
2.5	(SEQ ID NO: 6);	
35		
	CTANAVTGPC RASPPRWYFD VERNSCHNFI YGGCRGNENS YRSEE	150
	ACMLRC	156

SEQ ID NO: 7

	THOF CLUSKUVGEC FASMERWAYN VTDGSCQLEV TGGCDGNSMN	50
	YLTKEECLKK CATUTENATG DLATSRNAAD SSVPSAFFFÇ DSEDHSSOMF	75
5	MYEEYCTAMA VTGPORASEP RWYFDVERNS ONNFIYOGOR GMWWSYESEE	125
	ACMURCERQ	159
	SEQ ID NO: 3.;	
	CLVSKVVGRC RASMFRWWYN VTDGSCQLFV YGGCEGNSNN	50
10	YLTKEECLKK CATTENATG DLATSRNAAD SSVPSAPRRQ DSEDHSSDMF	100
	NYEETCTANA VTGTCRASFO RWYFDVERNS CNNFIVOGCR GNKNSYRGEE	150
	ACMLRC	156
	(SEQ ID NO: 50-)	
15	ADREPSIHOF CLUSHUNGRO FASMPRWWYN UTOGSOQLFU YGGODGNSMN	25
	YLTKEECLKK CATVTENATG CLATSFNAAD SSVPSAFRRQ DSEDHSSOMF	<u>. 2</u>
	NYEEYCTANA VTGFCRASFP RWYFDVERNS CHNFIYGGCR GHWNSYRSEE	125
	ACMLROFRQQ ENPPLPLGSK VVVLAGAVS	179
	(SEQ ID NO: 1); and	
20		
	ADRERSIHDE CLUSKUVGRC RASMPRWWYN UTDGSCQLEV YGGCDGNSNN	50
	YLTKEECLKK CATATENATS BLATSRNAAD SSVPSAFRRQ ES	92
	(SEQ ID NO: 8).	

- 25 2. A protein as in claim 1, wherein said protein is glycosylated, or contains at least one intra-chain cysteine-cysteine disulfide bond, or is both glycosylated and contains at least one intra-chain cysteine-cysteine disulfide bond.
- 3. A pharmaceutical composition for inhibiting serine protease activity.

  30 comprising a protein of claim 1 or claim 2 plus a pharmaceutically acceptable carrier.
- 35. A self-replicating protein expression vector containing a nucleic acid sequence which encodes for and is capable of expressing a protein of claim, or the containing an acid sequence which encodes for and is capable of expressing a protein of claim, or the containing an acid sequence which encodes for and is capable of expressing a protein of claim, or the containing an acid sequence which encodes for and is capable of expressing a protein of claim.

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6. A method for inhibiting serine protease activity comprising contacting serine protease with an effective amount of at least one protein of claim 1 or claim 2.

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7. A method for treating a condition of brain edema, spinal cord edema, multiple sclerosis, ischemia, perioperative blood loss, sepsis, septic shock, fibrosis, disease associated with pathologic blood coagulation or clotting, polytrauma, stroke, cerebral or subarachnoid hemorrhage, inflamation of the brain, inflamation of the spinal cord, cerebral infection, cerebral granulomatosis, spinal infection, spinal granulomatosis, open heart surgery, gastric cancer, cervical cancer, or prevention of metastasis comprising administering to a subject having such a condition an effective amount of the protein of claim 1 or claim 2.

15

8. The method of Claim 7 wherein said condition is brain edema, spinal cord edema, multiple sclerosis, ischemia, perioperative blood loss, sepsis, septic shock, fibrosis, disease associated with pathologic blood coagulation or clotting, stroke, cerebral or subarachnoid hemorrhage, inflamation of the brain, inflamation of the spinal cord, cerebral infection, cerebral granulomatosis, spinal infection, spinal granulomatosis, or open heart surgery.

25

30

20

- 9. The method of Claim 7 wherein said condition is gastric cancer, cervical cancer, or prevention of metastasis.
- 10. A method for the preparation of a medicament for the treatment of brain edema, spinal cord edema, multiple sclerosis, ischemia, perioperative blood loss, sepsis, septic shock, fibrosis, disease associated with pathologic blood coagulation or clotting, stroke, cerebral or subarachnoid hemorrhage, inflamation of the brain, inflamation of the spinal cord, cerebral infection, cerebral granulomatosis, spinal infection, spinal granulomatosis, open heart surgery, gastric cancer, cervical cancer, or prevention of metastasis.
- 11. A method for preparing a protien of claim 1 or claim 2 using recombinant DNA technology.

# FIGURE 1

ORF		F S P				50 16
R35464		ACCGAGAACG R E R			TGGTGTCGAA	
JRE	W E A D	RER	2 1 H	D F C L	VSK	3 3
	GGTGGTGGGC	AGATTCCGGG	CCTCCATGCC	TAGGTGGTGG	TACAATGTCA	150
ORF	v v G	R E R A	S M P	R W W	Y N V T	50
R35464	CTGACGGATC	CTGCCAGCTG	TTTGTGTATG	GGGGCTGTGA	CGGAAACAGC	200
ORF		C Q L			G N S	66
R35464	AATAATTACC	TGACCAAGGA	GGAGTGCCTC	AAGAAATGTG	CCACTGTCAC	250
ORF	и и у г	T K E	E C L	K K C A	T V T	83
R35464	AGAGAATGCC	ACGGGTGACC	TGGCCACCAG	CAGGAATGCA	GCGGATTCCT	300
ORF	E N A	T G D L	A T S	RNA	A D S S	100
R35464	CTGTCCCAAG	TGCTCCCAGA			ACTTCAGCGA	350
ORF	V P S	A P R	R Q D S	• R P	L Q R	116
R35464	TATGTTTCAA	NTATTGNAAG	AATAATTGCA	CCGNCAACGN	ATT	393
ORF	Y V S *	I * R	I I A	P * T *		130
KEY						
		cid sequenc			D NO: 12)	
	NE MINARA (	INAN WARKING	Frame Trac		O TO MO. 13'	

### FIGURE 2

R7 <b>4593</b> ORF		P D Q G		Q E M	C H C H	17
R74593 ORF	ACAGAGAATG R E C	CCACGGGTGA H G *	CCTGGCCACC P G H Q	AGCAGGAATG Q E C	CAGCGGATTC S G F	100 33
R74593 ORF	CTCTGTCCCA L C P K			TCTGAAGACC S E D H	ACTCCAGCGA S S D	150 50
R74593 ORF	TATGTTCAAC M F N	TATGAAGAAT Y E E Y			ACTGGGCCTT T G P C	200 67
R74593 ORF		CTTCCCACGC F P R	TGGTACTTTG W Y F D		GAACTCCTGC N S C	250 83
R74593 ORF		TCTATGGAGG Y G G		AATAAGAACA N K N S	GCTACCGCTC Y R S	300 100
R74593 ORF	TGAGGAGGCC E E A	TGCATGCTCC C M L R		CCAGCAGGAG Q Q E	AATCCTCCCC N P P L	350 117
R74593 ORF		CTCAAAGGTG S K V			TTCGTGATGG S * W	400 133
R74593 ORF		TTTCCTGGGG S W G		GTCTTACTGA V L L I	TTCCGGGTGG P G G	450 150
R74593 ORF	CAAGGAGGAA K E E		GCCCTGCGGA P A X		CTTCGGAGAT L R R *	500 167
R74593 ORF	GACAAGGGNT Q G					510 169

KEY

R74593 = Nucleic acid sequence of EST R74593 (SEQ ID NO: 14)
ORF = EST R74593 Open Reading Frame Translation (SEQ ID NO: 15)

# FIGURE 3

R35464	GGCCGGGTCGT	TTCTCGCCTG	GCTGGGA-TC	GCTGCTCCTC	TOTGGGGTCC 50
N397 <b>98</b>			TGGGANTC	GCTGCTCCTC	TOTGGGGTCC 28
H94519	GCNGCG-CGT	TNNTCGCNT-	GCTGGGA-TC	GCTGCACCTC	TOTGGGGTCG 47
874593 corr.					
Consensus	GGCCG <b>GGT</b> CGT	TTCTCGCCTG	GOTGGGA-TO	GCTGCTCCTC	TOTGGGGTCC 50
Translation	A G S F	L A W	L G S	L L L	S G V -3
1141131401011	r 3 3 :	<b>□</b> Λ η	2 3 3		3 3 V -3
D35464	TOCOCCOCC	ACCCACAACC	CACCATCCAC	CACTECECC	macamana 100
R35464					TGGTGTCGAA 100
N39798		ACCGAGAACG			TGGTGTCGAA 77
H94519	NGG-CGGCCG	ACCGAGAACG	CAGCATCCAC	GACTTCTGCC	TGGTGTCGAA 36
R74593 corr.					
Consensus		ACCGAGAACG		GACTTCTGCC	TGGTGTCGAA 99
Translation	L A A D	R E R	S I H	D E C L	Y S K 15
R35464	GGTGGTGGGC	AGATTCCGGG	CCTCCATGCC	TAGGTGGTGG	TACAATGTCA 150
N39798	GGTGGTGGGC	AGATGCCGGG	CCTCCATGCC	TAGGTGGTGG	TACAATGTCA 127
H94519	GGTGGTGGGC	AGATGCCGGG	CCTCCATGCC	TAGGTGGTGG	TACAATGTCA 146
R74593 corr.					
Consensus	GGTGGTGGGC	AGATGCCGGG	CCTCCATGCC	TAGGTGGTGG	TACAATGTCA 149
Translation	y y g	R C R A	SMP	B W W	Y N Y T 32
					<u>_</u>
R35464	CTCACCCATC	CTGCCAGCTG	TTTGTGTATG	SCCCCTCTCX	CGGAAACAGC 200
N39798		CTGCCAGCTG			
	-		-		
H94519	CTGACGGATC	CTGCCAGCTG	FILGIGIATG	GGGGCTGTGA	_
R74593 corr.					GC 2
Consensus		CTGCCAGCTG			
Translation	D G S	COL	E Y X G	G C D	G N S 48
R35464	AATAATTACC	TGACCAAGGA	GGAGTGCCTC	AAGAAATGTG	CCACTGTCAC 250
N39798	AATAATTACC	TGACCAAGGA	GGAGTGCCTC	AAGAAATGTG	CCACTGTCAC 227
H9 <b>4519</b>	AAT <b>AATTA</b> CC	TGACCAAGGA	GGAGTGCCTC	AAGAAATGTG	CCACTGTCAC 246
R74593 corr.	AATAATTACC	TGACCAAGGA	GGAGTGCCTC	AAGAAATGTG	CCACTGTCAC 52
Consensus	AATAATTACC	TGACCAAGGA	GGAGTGCCTC	AAGAAATGTG	CCACTGTCAC 249
Translation	NNYL	T K E	E C L	K K C A	T V T 65
R35464	AGAGAATGCC	ACGGGTGACC	TGGCCACCAG	CAGGAATGCA	GCGGATTCCT 300
N39798		ACGGGTGACC			
H94519		ACGGGTGACC			
R74593 corr.					
		ACGGGTGACC			
Consensus		ACGGGTGACC			
rranslation	E N A	1 6 2 7	A 1 5	RNA	A D S S 32
R35464	CTGTCCCAAG	TGCTCCCAGA	AGGCAGGATT	CTTGAAGACC	ACTTCAGCGA 350
N397 <b>98</b>	CTGTCCCAAG	TGCTCCCAGA	AGGCAGGATT	CT-GAAGACC	ACTCCAGCGA 326
H9 <b>4519</b>	CTGTCCCAAG	TGCTCCCAGA	AGGCAGGATT	CT-GAAGACC	ACTCCAGCGA 345
					ACTCCAGCGA 151
Consensus	STGTCCCAAG	TGCTCCCAGA	AGGCAGGATT	TT-GAAGACC	ACTOCAGOGA 749
		= E : ***	18.4	A	
. 27 3 4	TATE TAN	ETA T. AND	MITACT BOA	JUGGCAACGG	AGTOACTOU.
494519	TATGTT-CAA	CTA-TG-AAG	AATACTGGCA	COSCOAACGO	ATTCACTGGG 191
					AGTCACTGGG 197
Conconsus	TATOLITUMA	CTA-13-MAG	ADDACT TOCK	CCCCCAACGC	AGTCACTGGG 234
Jonsen <b>su</b> s	.A.G.I-CAA	0.M-13-MMG	WYTHOTIARY	COSCIANCOC	MOTOWCIDED NIA
Trans. areses		-	-	N. AT A	- · · · · · · · · · · · · · · · · ·

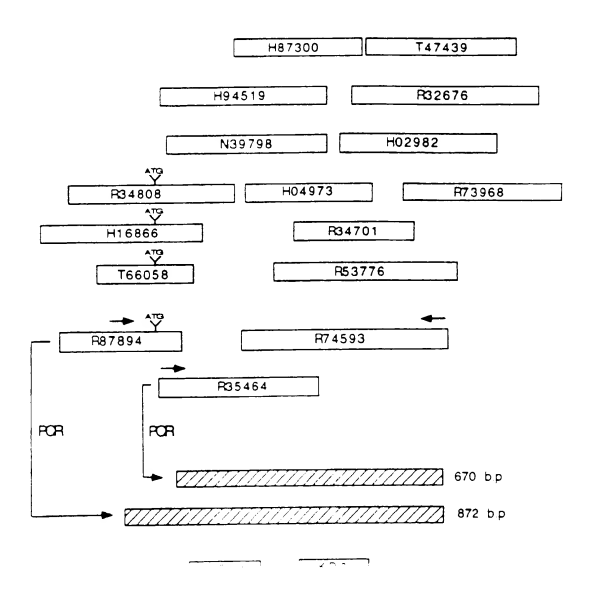
### FIGURE 3 (CON'T)

R35464						
N39798	CCTTGC-GTG	GAATCCTTTC	CCACGCTGGN	AATTTNGACG	TTGAGAAGGA	421
Н94519				ACTTT-GNCG		
R74593 corr.				ACTTT-GACG		
				ACTTT-GACG		
Consensus		S F	P R W Y			129
Translation	PCRA	3 :	r K W I	. 5	<b>L</b>	10,
R35464						
N39798	AC					423
H94519						
R74593 corr.	A CTCCTCC & A	TABCTTCATC	TATEGAGGCT	GCCGGGGCAA	TAAGAACAGO	293
				GCCGGGGCAA		490
Consensus		N F I	Y G G C	R G N	K N S	145
Translation	S C N	N F I	1 3 3 6	, G ,	K 14 3	143
R35464						
N39798						
H94519						
R74593 corr.	TACCECTCTE	AGGAGGCCTG	CATGCTCCGC	TGCTTCCGCC	AGCAGGAGAA	343
Consensus	TACCCCTCTG	AGGAGGCCTG	CATGCTCCGC	TGCTTCCGCC	AGCAGGAGAA	540
•••••		E A C	M L R	C F R Q	Q E N	162
Translation	YRSE	E A C	n b K	C : K V	2 2	
R35464						
N39798						
Н94519						
R74593 corr.	TOCTOCOCTG	CCCCTTGGCT	CAAAGGTGGT	GGTTCTGGCC	GGGGCTGTTT	393
				GGTTCTGGCC		590
Consensus		P L G S	K V V	V L A		179
Translation	P P L	P L G 3	K V V	V L A	G A V J	
R35464						
พ39798						
н94519						
R74593 corr.	CGTGATGGTG	TTGATCCTTT	TCCTGGGGAG	CNTCCATGGT	CTTACTGATT	443
	CGTGATGGTG			CNTCCATGGT		640
Consensus	* W C	* S F	S W G A		L L I	195
Translation	- w C	- 3 F	3 W G A	3 H V		.,,
R35464						
N39798						
H94519						
R74593 corr.	CCCCCTCCCA	AGGAGGAACC	AGGAGCGTGC	CCTGCGGANC	GTCTGGAGCT	493
				CCTGCGGANC		690
Consensus	CCGGGIGGCA	AGGAGGAACC	7007000100	D. A. T. B.	7 6 7	
Translation	P G G K	E E P	G A C	PA-R		
R35464						
N39798						
H94519						511
R74593 corr.						708
Consensus	TCGGAGATGA					
Translation	R R *	Q G				217
KEY			am nac464 :	CEO TO NO :	121	
R35464 = Nucl	eic acid se	dneuce of E	51 KJ5464 (	SEQ ID NO.:	171	
N39798 = Nucl	eic acid se	quence of E	ST N39/98 (	SEQ ID NO.:	1/)	
H94519 - Nucl	eic acid se	quence of E	ST H94519 (	SEQ ID NO.:	10)	
R74593 COFF	= Corrected	version of	(SEQ ID NO	),: 14) G at	b.p. 114	

R74593 corr. = Corrected version of (SEQ ID NO.: 14) G at b.p. 114 Consensus = Nucelic acid sequence for human bikunin (SEQ ID NO.: 9) Translation = Amino acid Translation of Consensus (SEQ ID NO.: 10)

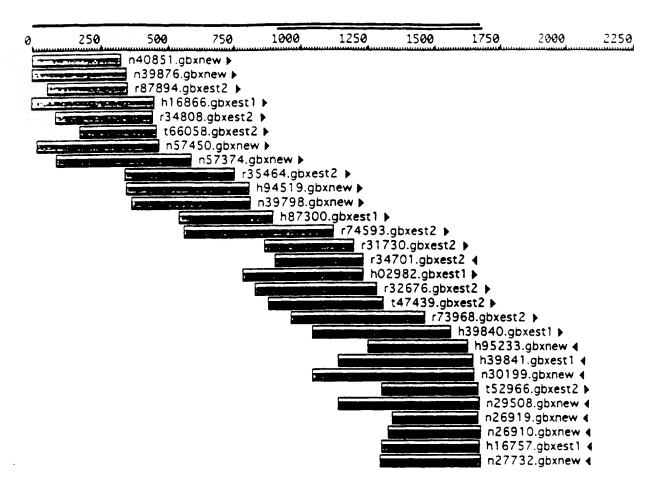
Figure 4 A.

Schematic depicting the overlap of ESTs bearing homology to the cDNA sequence encoding placental bikunin



Base pairs

Figure 4B



# Figure 4C

•	:				50
Bikunin	GCGA	compagadas	TTGGGAGGTG	TAGCGCGGCT	CTGAACGCGT
N40851	GCGA	cctccscscs	TTGGGAGGTG	TAGCGCGGCT	CTGAACGCGT
N39876	GCGA	corcososos	TTGGGAGGTG	TAGCGCGGCT	CTGAACGCGT
R87894					
H16866	L GGCGA	corocacaca	TTGGGAGGTG	TAGCGCG.CT	CTGAACGGGN
R34808					
766058					
N57450				TAGCGCGGCT	CTGAACGCNA
พร7374					
R35464					
H94519					
N39798					
H87300					
R74593					• • • • • • • • • •
R31730					
R34701					
H02982					
R32676					
T47439			• • • • • • • • •		
R73968					
H39840					
H95233	• • • • • • • • • •				
H39 <b>84</b> 1					• • • • • • • • • • • • • • • • • • • •
N30199	• • • • • • • • • • • • • • • • • • • •	· · · · · · · · · · · · · · · ·			
T52966			• • • • • • • • • • • • • • • • • • • •		
N29508		***************************************	• • • • • • • • • • • • • • • • • • • •		
N26919					
N26910					
H16757		• • • • • • • • •			
N27732					

3		100	<i>'</i>		
	51				100
Bikauiu	GNA GGGCCG	TIGAGIGICG	CAGGCGGCGA	GGGCGCGAGT	GAGGAGCAGA
N40851	NGAGNGGCCG	TTGAGTGTCG	CAGGCGGCGA	GGGCGCGAGT	GAGGAGCAGA
N39876	GCA.GGGCCG	TIGAGIGICG	CAGGCGGCGA	GGGCGCGAGT	GAGGAGCAGA
987894		TIGAGIGING	NAGGCGGCGA	GGGCGCGAGT	GAGGAGCAGA
H16866	ANGGGCCG	TTGAGTGTCG	CAGGCGGC . A	GGGCN.GAGT	GAGGAGCAGA
R34808					GAGGAGCAGA
T66058					
N57450	GAAGNGGCCG	TTGAGTGTCG	CAGGCGGCGA	GGGGGGAGT	GAGGAGCAGA
N57374					AGA
R35464			• • • • • • • • • • • • • • • • • • • •		
H94519					
N39798					
H87300					
R74593					
R31730					
R34701					
H02982					
R32676					
747439					
R73968					
H39840					
H95233					
H39841					
N30199					
T52966				• • • • • • • • • • • • • • • • • • • •	
N2 9508					
N26919					
N26910					
H16757					
N27732					

# Figure 4C (Con't)

	101				150
Bikunin	CCCAGGCATC	GCGCGCCGAG	AAGNC GGGC	GTCCCCACAC	TSAAGGTCCG
N40851	CCCAGGCATC	GCGCGCCGAG	AAGNC.GGGC	GTCCCCACAC	TSAAGGTCCG
N39876	CCCAGGCATC	GCGCGCCGAG	AAGNC . GGGC	NTCCCCACAC	TGAAGGTCCG
R87894	CCCAGGCATC	GCGCGCCGAG	AAGGCCGGGC	GTCCCCACAC	TGAAGGTCCG
H16866	CCCAGGCATC	GCGCGCGAG	AAGNC.GGGC	GTCCCCACAC	TGAAGGTCCG
R348C8	JCCAGGCATC	GCGCGCCGAG	AAGNC.GGGC	GTCCCCACAC	TGAAGGTCCG
T66058					
N57450	CCCAGGCATC	GCGCGCGAG	AAGNC.GGGC	GTCCCCACAC	TGAAGGTCCG
N57374	CCCAGGCATC	GCGCGCCGAG	AAGNC, GGGC	GTCCCCACAC	TGAAGGTCCG
R35464					
H94519					
N39798					
H87300					
R74593					
R31730					
R34@01		• • • • • • • • •			
HC2982					
R32676			• • • • • • • • •		• • • • • • • • •
T47439		• • • • • • • •			• • • • • • • • •
R73968			• • • • • • • • • •	• • • • • • • • • •	
839840	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • •		• • • • • • • • • • • • • • • • • • • •
H95233		• • • • • • • • • •		• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
H39841	· · · · · · · · · · · · · · · · · · ·		• • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	
N30199			• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • •
752966			• • • • • • • • • •	• • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
N29508				• • • • • • • • •	
N26919				• • • • • • • • •	
N26910					
H16757				• • • • • • • • •	
N27732					

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3		(0020	,		
	151				200
Bikunin	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	GGCGGACCCT	CCCGGAGCGT
N40851	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	GGCGGACCCT	CCCGGAGCGT
N39876	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	GGCGGACCCT	CCCGGAGCGT
R87894	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	GGCGGACCCT	CCCGGAGCGT
H16866	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	GGCGGACG.T	CCCGGAGCN.
R34808	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	GGCGGACCCT	CCCGGAGCGT
T66058				GGACCCT	CCCGGAGCGT
N57450	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	GGCGGACCCT	CCCGGAGCGT
N57374	GAAAGGCGAC	TTCCGGGGGC	TTTGGCACCT	GGCGGACCCT	CCCGGAGCGT
R35464					
H94519					
N39798					
H87300	• • • • • • • • • • • • • • • • • • • •				
R74593	• • • • • • • • • • • •				
R31730					
R34701					
H02982					
R32676					
T47439					
R73968		)			
H39840					
H95233					
H39841					
N30199	• • • • • • • • • • • • • • • • • • • •				
T52966	• • • • • • • • • •				
N29508	• • • • • • • • •				
N26919	• • • • • • • • • • • • • • • • • • • •				
N26910	• • • • • • • • • • • •				
H16757					
N27732					

#### Figure 4C (Con't) 201 250 Bikunin CGGCACCTGA ACGCGAGGCG CTCCATTGCG CGTCCGTTTG .AGGGGCTTC N40851 CGGCACCTGA ACGCGAGGCG CTCCATTGCG CGTGCGTNTG , AGGGGCTTC N39876 CGGCACCTGA ACCCGAGGCC CTCCATTGCC CGTGCGTTTG .AGGGGCTTC RBTB94 CGGCACCTGA ACGCGAGGGG CTCCATTGCG CGTGCGTTTG .AGGGGCTTC HIEROROPA ACCORDANCE CONTROL CONTROL ACCORDANCE CONTROL CONTRO RB4808 CGGCACCTGA ACGCGAGGCG CTCCATTGCG CGTGCGTNTG GAGGGGCTTC TE6058 CGGCACCTGA ACGCGAGGC, CTCCATTGCG, GTGCGTGTG NAGGGGGCTTC NER450 CGGCACCTGA ACGCGAGGCG CTCCATTGCG CGTGCGTTTG .AGGGGCTTC NS1374 CGGCACCTGA ACGCGAGGC, CTCCATTGC, CGTGCGTTNG , AGGGGCTTC R35464 ..... 894519 ...... N39798 H87300 ..... 274593 ..... R31730 ...... 234701 H02982 R32676 ..... 747439 ..... 373968 ......... 339840 H95233 N30199 ...... T52966 ..... N29508 ..... N26919 ..... N2 6910 ...... H16757 ...... N27732 .....

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	251				300
Bikaniu	CCGCACCT G	ATCGCGAGAC	CCCAACGGCT	SGTGG CSTC	GC TG CGCG
N40851	CCGCACCT.G	ATCGCGAGAC	CCCAACGGCT	GGTGG.CGTC	SCCTS.CSCS
N39876	CCGCACCT.G	ATCGCGAGAC	CCCAACGGCT	SGTGG.CGTC	SCCTS.CSCS
R87894	CCGCACCT.G	ATCGCGAGAC	CCCAACGGCT	GGTNG.CGTC	GC.TN.CGCG
H16866	CCGCACCT.G	ATCGCGAGAC	CCCAACGGCT	GGTNG.CGTC	GC.TGGCGCG
R34808	CCGCACCT.G	ATCGCGAGAC	CCCAACGGCT	GGTGGGCGTI	GC.TG.CGCS
T66058	CCGCACCT.G	ATCGCGAGAC	CCCAACGGCT	GGTGG.CGTC	GC.TG.CGCG
N57450	CCGCACCT.G	ATCGCGAGAC	CCCAACGGCT	SGTGG.CGTC	GCCTS.CGCS
N57374	CCGGAACTTG	ATCGCGAGAC	CCCAACGGCT	GGTGG.CGTC	GC.TG.CGCG
R35464					
H94519			· · · · · · · · · · · · · · ·		
N39798					
H87300					
R74593					
R31730					
R34701					
H02982					
R32676					
T47439					
R-3968					
H39840					
H95233					
H39841					
830199					
T52966					
N29508					
N26919					
N26910					
H16757					
N27732					

	30:	
_		350
Bikunin	TO TODGETS AGET GGCCA TOGGGGCANT GTTGC GGGC T GAGGC	ಽಽ
N40851	TO TEGGETS AGET, GENEA TETES	
439876	TO TOGGOTS AGOT GGOCA ISGOSCACT, SITGOSGNGC TIGAGGO.	. 3
387894	TO TOGGOTG AGETTGGCCA TGGCGCANT, GTTNC, GGGC T.NAGGC.	. 33
H16866	TICTOGGOTG AGCT.GGCCA TGGCGCANT, GTTGC.GNGC T.GAGGC.	SS
834808	TOTTOGGOTG AGCTGGGGCA TGGCGCANTT GTTGC.GGGC T.GAGGC.	GG
766058	TO TOGGOTG AGOT.GGCCA TGGCGCANT. GTTGC.GNGC T.GAGGC.	GG
N57450	TO TOGGOTG AGOT, GGCCA TGGCGCANT, GGTGC, GGGC TTGAGGC.	
N57374	TESTEGGETG AGET.GGCCA TGGCGCANT. GGTGCCGNGC T.GAGGCC	CG
R35464		GG
H94519		
N39798		
n87300		
R74593		
P31730		
P34701		
902982		
32616		
T47439		
373968		
H39840		
H95233	***************************************	
H39841		
N30199		
T52966		
N2 95 08		
N26919		
N26910		
H16757	***************************************	
V27732	********	

	351				400
Bikunin	AC GG CG	TTTCTCS	co tactass	A TOGOT SC	T COTOTOT
R87894	ACG.				
H16866	ACCGNCGT	TITTCTTCS.	CCTTSCTGGG	ATTOGCTTSC	TTCCTNTCTC
234806	ACGCGGNCG.	LITITITICSN	CCTTGCTGGG	ATTCG.TTG.	TENCTOTOEN
T66058	CGGNCG.	.TTTTCTCS.	cc.tsctss	A.TOGOT.GO	T.COTOTOT.
N57450	ANN.NGCCG.	TITCTCS.	CC.TGCTGGG	A.TOGOT.GO	T.COTOTOT.
N57374	AGGGCCGG	ITTCTCG.	CCTTGCTGGG	A.TCGCT.GC	T.CCTCTCTC
R35464	GTCG.	ITTCTCG.	CCTGGCTGGG	A.TCGCT.GC	T.CCTCTCT.
H94519	GCNGCGCG.	ITNNTCG.	CN.TGCTGGG	A.TOGOT.GO	A.COTOTOT.
N39798			CTGGG	ANTOGOT.GO	T.COTOTOT.
H87300					
274593					
R31730					
R34701					
H02982					
R32676					
147439					
R73968					
H39840					
H95233					
H39841					
N30199					
T52966	• • • • • • • • • • •				
N29508					
N26919					
N26910					
816757					
N27732					

	401				450
Bikunin	GGGG TEETS	G COGCOGA	CCGA GAACG	CA SCA TCC	ACGACTT CT
H16866	GGGGTTCCTG	GG:CGGCCGA	CCGA, SAACG	CA.SCA.TCC	AAGAATTTT
R34808	GGGGTTC.TS	GGGNGGCCGA	NCGA, GAACG	CAAGCA.TTC	ACGA.TTT
166058	GGGG. TCCTS	S CGGCCGA	CCGA.GAACG	CALGCALTCC	ACGANTT.CT
N57450	GGGG.TCCTG	G CGGCCGA	COGA,GAACG	CALSCALTCC	ACGACTT.CT
N57374	GGGG, TCCTG	GCGGCCGA	NCGAAGAANG	CA.SCAATCC	ANGAATINCT
R35464	GGGG.TCCTG	G.CCGGCCGA	CCGA,GAACG	CA.GCA.TCC	ACGACTT.CT
H94519	GGGG.TCSNG	GCGGCCGA	CCGA.GAACG	CA.GCA.TCC	ACGACTT.CT
N39798	GGGG. TCCTG	GCGGCCGA	CCGA.GAACG	CA.SCA.TCC	ACGACTT.CT
H87300					
274593					
R31730		• • • • • • • • • • • • • • • • • • • •			
R34701					
H02982					
R32676					
T47439					
273968			**************************************		
H39840	• • • • • • • • • • • • • • • • • • • •			· · · · · · · · · · · ·	
H95233	• • • • • • • • • • • • • • • • • • • •				
839841	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • •		
N30199				• • • • • • • • • •	
T52966	• • • • • • • • • • • • • • • • • • • •				
N29508	• • • • • • • • • • • • • • • • • • • •				
N26919	• • • • • • • • • • • • • • • • • • • •				
N26910	• • • • • • • • • • • • • • • • • • • •		• * • • • • • •		
H16757	• • • • • • • • • • • • • • • • • • • •				
N27732					

	451				500
Bixunin	GCCTGGTGT	CS.VAGGT GG	TGGGCAGATG	cesss cere	CATGCCTA G
H16866	SCC				
766058	TCCTGGTGTT	CGAAGG			
N57450	GCCTSSTST.	CGAAGGT.GG	TGGGCAG		
N57374	GCCTGGTGTT	CGAAAGTTGG	TGGGCANATT	CCGGGGCCTT	CATGNCTAAG
R35464	GCCTGGTGT.	CGAAGGT.GG	TGGGCAGATT	ccsgg.cctc	CATGCCTA.S
H94519	GCCTGSTGT.	CGAAGGT.GG	TGGGCAGATG	CCGGG.CCTC	CATGCCTA.G
N39798	GCCTGGTGT.	CGAAGGT.GG	TGGGCAGATG	ccsss.cctc	CATGCCTA. G
H87300					
R74593					
R31730					
R34701					
H02982					
R32676	• • • • • • • • • • • • • • • • • • • •				
T47439					
R73968					
839840					
H95233					
839841					
N30199					
T52966					
N29508	• • • • • • • • •				
N26919					
N26910					
H16757					
N27732					

Figure 4C (Con't) 550 S TOST OUT ACABTGTCAC TGACGGATCC TSCCAGCTGT TTGTGT ATG Bikunin 857374 GTTGGTTGGT ANAATGTNAA TTAANGATTC TTGCAACTGT TTGTGTNATT R35464 G. TOGT. GGT ACAATGTCAC TGACGGATCC TGCCAGCTGT TTGTGT.ATG #94519 SUTGGT.GGT ACAATGTCAC TGACGGATCC TGCCAGCTGT TTGTGT.ATG 839798 G.TGGT.GGT ACAATGTCAC TGACGGATCC TGCCAGCTGT TTGTGT.ATG R31730 ..... R32676 ..... ...... #3984C ....... ..... ..... ..... ...... N30199 N26919 N26910 #16757 ....... N27732 551 600 GGGGCTGTGA CGGAAACA GCAATAATTA CCTGACCAAG GA GGAGTGC Bigunin NSTB74 GGGGCTNTTA AACGGAAANA .CAATAATNA CCTGACCAAA GAAGNAAT... R35464 GGGGGTGTGA ...CGGAAACA GCAATAATTA CCTGACCAAG GA.GGAGTGC H94519 GGGGTTGTGA ... CGGAAACA GCAATAATTA CCTGACCAAG GA.GGAGTGC N39798 GGGGCTGTGA ...CGGAAACA GCAATAATTA CCTGACCAAG GA.GGAGTGC H97300 SATTOSCAC AGGGGAAACA GCAATAATTA COTGACCAAG GALGGAGING P74593 ..... GGAGTGG CATAATTA CCTGACCAAG GA GGAGTGC 931730 F34101 H02982 P32676 T4~439 273968 H39840 495233 -39841 V20199

715 E N27732

Fiç	jure 4	C	(Con't	)		
	601					653
Bikunin	CTCAAGAA	LAT.	GTGCCACTGT	CACAGAGAAT	GCCACGGGTG	ACCTGGCCAG
R35464	STCAAGAA	TA	GTGCCACTGT	CACAGAGAAT	GCCACGGGTG	ACCTGGCCAC
H94519	CTCAAGAA	TA	GTGCCACTGT	CACAGAGAAT	GCCACGGGTG	ACCTGGCCAG
N39798	CTCAAGAA	AT	GTGCCACTGT		GCCACGGGTG	
H87300	CTCAAGAA	AT	GINCCACIST		GCCACGGGTG	
R74593	CTCAAGAA	AT	GTGCCACTGT		GCCACGGGTS	
R31730						
R34701	· • • • • • • •					
HC2982						
R32676						
T47439						
R73968						
H39840						
H95233						
H39841	, , , , , , , ,					
N30199						
T52966						
N295C8						• • • • • • • • •
N26919						• • • • • • • • •
N26910						
H16757						
N27732		• •				
		• •				• • • • • • • • • •
	651					700
Bikunin		AT (	SCACCGGATT	CCTCTCTCC	AAGTGCTCCC	
R35464					AAGTGCTCCC	
H94519					AAGTGCTCCC	
N39798					AAGTGCTCCC	
H87300						
R74593					AAGT.CTCCC	
831730						
R34701						
802982			• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	
P32676						
147439				• • • • • • • •	• • • • • • • • • • • • • • • • • • • •	
R73968	· · · · · · · · ·				• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
H39840	• • • • • • •				• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •
H95233						
H39841						
N30199						• • • • • • • • • • • • • • • • • • • •
752966						
N29508						
N26919	· · · · · · · · ·					
N26910						
H16757						
N:7772						

Figure 4C (Con't) 150 BEXURER ATTOT GAAG ACCACTCCAG CGATATGTT CAACTAT G AAGAATACTG R35464 ATTCTTGAAG ACCACTTCAG CGATATGTTT CAANTATTGN AAGAATAATT H94519 ATTOT.GAAG ACCACTCCAG CGATATGTT, CAACTAT..G AAGAATACTG N39798 ATTCT.GAAG ACCACTCCAG CGATATGTT. CAACTAT...G AAGAATACTG H87300 ATTCT.GAAG ACCACTCCAG CGATATGTT, CAACTAT... AAGAATACTG R74593 ATTCT.GAAG ACCACTCCAG CGATATGTT. CAACTAT..G AAGAATACTG 931730 R34701 ногова R32676 ..... T47439 ....... R73968 H39840 ...... E95233 ....... 839841 N30199 752966 N26919 N26910 ...... H16757 N27732 ..... 751 800 CACCGCCAA CGCAGT CAC TGGGCC TTG CCGTG CAT CCTT CCCAC Bikunin R35464 GCACCGNCAA CGNATT H94519 GCACCGCCAA CGCATT.CAC TGGGCC..TG C.GTG.CAT. CCTT.CCCAC N39798 .CACCECCAA CECAGT.CAC TEGESCOTTE C.ETEGAAT. COTTTECCAC LIACOSCOAN OGGAGINGAC IGGGGG.TIG G.GIGGGAIN COIT.COCAC H87300 R14593 .CACCOCCAA CGCAGT.CAC TGGGCC.TTG CCGTG.CAT. CCTT.CCCAC 934701 H02982 R32676 T47439 ..... 273968 H39840 495233 39841 430199 752366 N29508

£ 1	gure 40	(Con't	; )		
	801				850
Bikunin	GCTGGTACT	T GACGTGGA	GA GGAACTO	CTS CAATAA	CTTCATCTAT
H94519	GCTGGTACT	T.GNCGT			
N39798	GCTGGNAATT	TNGACGTTGA	GAAGGAAC		
H87300	GCTNGTACTT	T.GACGTGGA	GA.SGAACTS	CTGGCAATAA	CTTCATCTAT
R74593	GCTGGTACT	T.GACGTGGA	GA. GGAACTO	CTG.CAATAA	CTTCATCTAT
R31730					
R34701					
HC2982	· · · · · · · · · · · · ·	GA	GA.GGAACTC	CTG.CAATAA	CTTCATCTAT
R32676					ATTCGGAA
T47439					
R73968					
H39840					
H95233					
H39841					
N30199			• • • • • • • • • • • • • • • • • • • •		
T52966	• • • • • • • • • • • • • • • • • • • •				
N29508					
N26919			• • • • • • • • • • • • • • • • • • • •		
N26910					
H16757	• • • • • • • • • • • • • • • • • • • •				
N27732					
	851				900
Bikunin	GGAGGCT GC	CGGGGCAAT	AAGAACAG C	TACCGCTC T	GAGGAGGCCT
H8730C	GGAGGCTTGC	CSGGGCAATN	AAGAACAGNT	TACCGCTCTT	TAGGAGGCCT
R74593	GGAGGCT.GC	CGGGGCAAT.	AAGAACAG.C	TACCGCTC.T	GAGGAGGCCT
R31730				TACCGCTC.T	GAGGAGGCCT
R34701					
H02982	GGNGGCT.GC	CGGGG.AAT.	AAGAACA.NC	TACCGCTC.T	GAGGAGGCCT
R32676	CGAGGAGC	CGGGGCAAT.	AAGAACAG.C	TACCGCTC.T	GAGGAGGCCT
T47439	• • • • • • • • • • • • • • • • • • • •				NGGCCT
R73968					
H39840					
H95233	• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • •		
H39841	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •	• • • • • • • • • • •		
N30199	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •		
T52966	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
N29508	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •			
N26919		• • • • • • • • •			
N26910					
H16757	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
N2 7 7 32	• • • • • • • • • • • • • • • • • • • •				

Fi	gure 40	(Con't	: )		
	901				950
Bikunin	GCA TGCT	cactactic:	<b>.</b> sc		CA GCAGGA
H87300	.GCA.T				
R74593	.GCA.TGCT	cactactic	s sc		CA , GCAGGA
R31730	. GCA . TGCT	coctocttc			
234701			GC		
HC2982	. GCG . TSCT:		: GCTGTGTGTT		GCA.GCAGGA
R32676	.GCA.TGCTC	cactactica	: sc		
T47439	TGCAGTGCTC	CGCTGCTTCC	: GC		. CA . GCAGGA
R73968					
H39840					
H95233					
H39841					
N30199					
T\$2966					
N29508					
N26919					
N26910					
H16757	• • • • • • • • • • • • • • • • • • • •				
N27732					
	951				1000
Bikunin		CCTGCCCCTT	GGCTCAAAGG	TGGTGGTTC	TGG CGGGGC
R74593	GAA . TOOTOO	CCTGCCCCTT	GGCTCAAAGG	TGGTGGTTC.	TGGCGGGGC
R31730	GAA. TOOTCO	CCTGCCCCTT	GGCTCAAAGG	TGGTGGTTC.	TGG.CGGGGC
R34701	AAANTCCTCC	CCTCCCCCTT	GGCTCAAAGG	TGGTGGTTCC	TGG.CGGGGC
H02982	GAA. TOOTCO	CCIGCCCCTT	GGCTCAAAGG	TGGTGGTTC.	TGG.CGGGGC
R32676	GAA. TCCTCC	CCTGCCCCTT	GGCTCAAAGG	TGGTGGTTC.	TGG. CGGGGC
T47439	GAA.TCCTCC	CCTGCCCCTT	GGCTCAAAGG	TGGTGGTTC.	TGG.CGGGGC
R73968					c <b>cccc</b> c
H39840		· · · · · · · · · · · · · · · · · · ·			
H95233	********				
H39841					
N30199	*				
T52966		0	*		
N29508	* * * * * * * * * * * * * * * * * * * *				
N26919		• • • • • • • • • • • • • • • • • • • •			
N26910		• • • • • • • • • • • • • • • • • • • •			
H16757					
N27732					

#### Figure 4C (Con't) 1001 1050 Bikunin TGTT CGTGA TGGTGTTGAT CC T CTTCC TGGG AGCCT CC ATGGTC R74593 TGTTTCGTGA TGGTGTTGAT CCTT..TTCC TGGGGAGCNT CC.ATGGTCT R31730 TGTT.CGTGA TGGTGTTGAT CC.T.CTTCC TGGGGAGCCT CC.ATGGTC. R34701 TGTT.CGTGA TGGTGTTGAT CCCTCCTTCC CGGG.AGCCT CCCATGGTCC H02982 TGTT.CGTGA TGGTGTTGAT CC.T.CTTCC TGGG.AGCCT CC.ATGGTN. R32676 TGTT.CGTGA TGGTGTTGAT CC.T.CTTCC TGGG.AGCCT CC.ATGGTC. T47439 TGTT.CGTGA TGGTGTTGAT CC.T.CTTCC TGGG.AGCCT CC.ATGGTC. R73968 TGTT.CGTGA TGGTGTTGAT CC.T.CTTCC TGGG.AGCCT CC.ATGGTC. H39840 ..... H95233 H39841 830199 T52966 ..... N29508 N26919 H16757 ..... N27732 1051 Bikunin TACC TGAT CCGGGTGGCA CGGAGG AAC C AGG AGCG TGCCCTGCGC R74593 TAC..TGATT CCGGGTGGCA AGGAGG.AAC C.AGG.AGCG TGCCCTGCGG R31730 TACC.TGAT. CCGGGTGGCA CGGAGGGAAC C.AGGGAGCG TGCCCTGCGC R34701 TACCOTGAT, COGGGTGGCA CGGAGG, AAC CCAGG, ANCG TGCCCTGCGC H02982 TACC.TGAT. CCGGGTNGCA CGGAGG.AAC C.AGGGAGCG TGCCCTGCGN R32676 TACC.TGAT. CCGGGTGGCA CGGAGG.AAC C.AGGGAGCG TGCCCTGCGC T47439 TACC.TGAT. CCGGGTNGCA CGGAGG.AAC C.AGG.AGCG TGCCCTGCGC R73968 TACC.TGAT. CCGGGTGGCA CGGAGG.AAC C.AGG.AGCG TGCCCTGCGC н95233 H39841 N30199 T52966 ..... N29508 ..... N26919 N26910 .....

Figure 4C (Con't) 1101 1150 Bikunin ACCS TOT G GAGCTCCGGA GATGACAAGG AGCAGCTGG TGAAGAAC P14593 ANCOLTCT.S SAGCTTCGGA GATGACAAGG GNT R31730 ACCG.TCTGG GAGCTCCGGA GATGACAAGG GAGCAGCTGG GTGAAGAAC. R34701 ACCC.TCT.G GAGCTCCGGA GATGACAAGG .AGCAGCTGG .TGAAGAAC. HO2982 ACCC.TCTNG GAGCTCCGGA GATGACAAGG .AGCAGCTGG .TGAAGAAC. R32676 ACCC.TCTGG GAGCTCCGGA GATGACAAGG GAGCAGCTGG .TGAAGAAC. T47439 ACCG.TCT.G GAGCTCCGGA GATGACAAGG .AGCAGCTGG .TGAAGAAC. R73968 ACCG.TCT.G GAGCTCCGGA GATGACAAGG .AGCAGCTGG .TGAAGAAC. H39840 ACCGGTCT.S GAGCTCCGGA GATGACAAGG .AGCAGCTGG .TGAAGAAC. Н95233 #39841 ...... N30199 ACCG.TCT.G GAGCTCCGGA GATNACAANG .AGCAGCTGN .TGAAGAACC T52966 ...... N29508 N26919 ..... N26910 ..... H16757 ..... N27732 1151 BIKUNIN ACATATOT C CTGT GACCO CCCTGT CGC C AAGAGG A CT GGGGAA R31730 ACATATGTTC CTGTTGACCG NCCTGTTCGC C.AAGAGG.A TTGGGGGAA. R34701 ACATATGT.C CTGT.GACCG CCCTGT.CGC C.AAGAGG.A CT.GGGGAA. H02982 ACATAIGT.C CTGT.GACCG NCCTGTTCGN C.AAGAGG.A CTNGGGGAAA R32676 ACATATGTTC CTGTTGACCG CCCTGTTCGC C.AAGAGGGA NTGGGGGAA. T47439 ACATATGT.C CTGT.GACCG CCCTGT.CGC C.AAGAGG.A CT.GGGGAA. R73968 ACATATGT.C CTGT.GACCG CCCTGT.CGC C.AAGAGG.A CT.GGGGAA. H39840 ACATATGT.C CTGT.GACCG CCCTGT.CGC C.AAGAGG.A CT.NGGGAA. H39841 .....C. CCCTGT.CGC CCAAAAGG.A CT.GGGGAA. N30199 ACATATGT.C CTGT.GACCG CCCTNT.CGC C.AAGAGG.A CT.GGGNAAA N29508 ......CC. CCCTNT.CGC C.AAGAGG.A CT.GGG.AA N26919 N26910 

Fi	gure 40	C (Con't	:)		
	1201				1250
Bikunin	GGGAGGGG		TOT GA GC	I ITITIT A	A TAGA GG
R31730	. GGGAGGGG				
R34701	. GGGAGGGG			: IIIIIIA/	A.TA
H02982	GGGGAGGGG				
R32676	GGGGAGGGG			CANTITITIT 1	ATTAGGAGGG
T47439	. GGGAGGGG		TGT.GA.GC1	TITITITI AA	A.TAGAGG
R73968	. GGGAGGGG	. AGACTAT.G.	TGT.GA.GCT	TITTTT AA	A.TAGAGG
H39840	. GGGAGGGG	. AGACTAT.G.	TGT . GA . GCT	LA. TITITIT	A.TAGAGG
H95233	• • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			
H39841		A AAACNAT.G.			A.TAGAGG
N30199	. GGGAGGNG	. AGACTAT.G.	TGT . AA . GCT	TITITI AA	A.TAGAGG
T52966	• • • • • • • • •				
N29508	. GGGAGGGG	. AGACTAG.	TGT.GA.GCT	TITITITAA	A.TAGAGG
N26919		• • • • • • • • • • • • • • • • • • • •		• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •
N26910					
H16757					
N27732	• • • • • • • •			• • • • • • • • • • • • • • • • • • • •	
	1251				1300
Bikunin	GATTGACTC		GT GATC A		GAGGTCTGTT
R32676	GNTTGANTTO		GTTGATCCAT		
T47439	GATTGACTC		GT.GATC.A.	TTAGGGCT	GAGGTCTNTT
R73968	GATTGACTC		GT.SATC.A.	TTAGGGCT	GAGGTCTGTT
H39840	GATTGACTO		GT.GATC.A.	TTAGGGCT	GAGGTCTGTT
Н95233	• • • • • • • • •		<b> </b>	TTAGGGCT	GAGGTCTGTT
H39841	GATTGACTC.		GT.GATC.A.		GAGGTCTGTT
N30199	GATTGACTC.		GT.GATC.A.	TTAGGGCT	GAGGTCTGTT
T52966			• • • • • • • • • • • • • • • • • • • •		
N29508	GATTGACTC.		GT.GATCNA.	TTAGGGCT	GAGGTCTGTT
N26919					
N26910		• • • • • • • • • • • • • • • • • • • •			
H16757					
N27732					
_	1301				1350
Bikunin		GTAGGACGGC	TUCTTOC TO	G TO TOGGA	GGGATGGG
747439		GTAGGACGA			
R73968		GTAGGACGGC			
H39840		GTAGGACGGC			
H95233		NTAGGACGGC			
H39841		GTAGGACGGC			
N30199		GTAGGACGGC			
T52966					
N29508		GTAGGACGGC			
N26919					
N26910					
d16757		• • • • • • • • • • • • • • • • • • • •			
N27732					

```
Figure 4C (Con't)
         1351
                                                         :400
Bikunin TTTG CTTTG G AMATOCTO T AGGAGGOT COTOCT CGC ATGG CC TS
 R73968 TITG.CITTG GGAAATCCTC TINGGAGGCT CCTCCTTCGC ATGGGCCTTG
 H39840 TTTG.CTTTG GAGAATCCTC T.ANGAGGCT CCTCCT.CSC ATGG.CC.TS
 H95233 TTTG.CTTTG G.AAATCCTC T.AGGAGGCT CCTCCT.CGC ATGG.CC.TG
 H39841 TITG.CITTG G.AAANCONC T.AGGAGGCT CCTCCT.CGC ATGG.CC.TG
 N30199 TTTG.CTTTG G.AAATCCTC T.AGGAGGCT CCTCCTTCGC ATGG.CC.TG
 T52966 TTTG.CTTTG G.AAATCCTC T.AGGAGGCT CCTCCT.CGC ATGG.CC.TG
 N29508 TTTG.CTTTG G.AAATCCTC T.AGGAGGCT CCTCCT.CGC ATGG.CC.TG
 N26910 .....CTTTT GNAAATCCTC T.AGGAGGCT CCTCCT.CGC ATGG.CC.TG
 H16757 TITESCETTE G. AMANCETE T. AGRAGGET CETCET, CGC ATGG.CC. TG
 N27732 TTTG.CTTTG G.AAATCCTC TTAGGAGGCT CCTCCT.CGC ATGG.CC.TG
        1401
                                                         1450
Bikunin CAST CT GG CAGCAG CCC CGAGTTGTTT CC TCGCTG ATC GATTTC
 273968 CAGT.CINGG CAGCANCOOC CGAGTTTTTT TCCTTCGCTG ATCCGATTTC
 H39840 CAGT.CT.GG CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ATC.GATTTC
H95233 CAGTTOTILG CAGCAGICOC CGAGTTGTTT LCC.TCGCTG ATC.GATTTC
H39841 CAGT.CT.GG CAGCAG.CCC CGAGTTGTTN .CC.TCGCTG ATC.GATNTC
N30199 CAGT.CT.GG CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ATC.GATTTC
 T52966 CAGT.CT.GG CAGCAG..CC CGAGTTGTTT .CC.TCGCTG ATC.GATTTC
N29508 CAGT.CT..G CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ATC.GATTTC
N26919 CAGT.CTTGG CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ANC.GATTTC
N26910 CAGT.CT..G CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ATCGGATTTC
H16757 CAGINOTIGG CAGCAGACCO CGAGITGITT .CC.TCGCTG ATC.GATITC
827732 CAGT.CT.GG CAGCAG.CCC CGAGTTGTTT .CC.TCGCTG ANC.GATTTC
        1451
                                                         1500
Bikunin ITT CCTCCA GGTAG AGT TTTC TTTG CTTATGTTGA ATTCCATTGC
R73968 ITTTCCTCCA GGTAAGAATT TTTCTTTT
H39840 TTT.CCTCCA GGTAG..AGT TTTC.TTTG, CTTATGTTGA ATTCCATTGC
H95233 TIT COTOCA GGTAG. AGT TITC.TITG. CITATGTTGA ATTCCATTGC
H39841 TTT.CCCCCA GGTAG..AGT TTTC.TTTG. CTTATGTTGA ANTCCATTGC
N30199 TIT.COTCCA GGTAG..AGT TITC.TITG. CITATGTTGA ATTCCATTGC
T52966 TIT.COTCCA GGTAG..AGT TITC.TITG. CITATGTTGA ATTCCATTGC
N29508 TIT.COTOCA GGTAG..AGT TITC.TITG. CTTATGTTGA ATTCCATTGC
N26919 TITLICONCCA GGTAGLLAGT TITCLITITG, CITATGTTGA ATTOCATTGC
N26910 TITLOCTOCA GGTAG. AGT TITC.TITG. CTTATGTTGA ATTCCATTGC
H16757 TTTACCCCCA GGTAG. .AGT TTTCCTTTGN CTTATGTTGA ATTCCATTGC
N27732 TTT.CCTCCA GGTAS. AST TTTC.TTTG. CTTATGTTGA ATTCCATTGC
```

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Figure 4C (Con't)
         1501
                                                         1550
BIRUNIN CICITIT OF CATCACAGAA GIGATOTIGG AATOGITTOT TITGITT OF
 H95233 CICTITI.CT CATCACAGAA GTGATGTIGG AATCGTTTCT TITGTTT.GT
 H39841 CTCTTTT.CT CATCACAGAA GTGATGTTGG AATCGTTTCT TTTGTTT.GT
 N30199 CTCTTTT.CT CATCACAGAA GTGATGTTGG AATCGTTTCT TTTGTTT.GT
 T52966 CTCTTTT.CT CATCACAGAA GTGATGTTGG AATCGTTTCT TTTGTTT.GT
 N29508 CTCTTTT.CT CATCACAGAA GTGATGTTGG AATCGTTTCT TTTGTTT.GT
 N26919 CTCTTTT.CN CATCACAGAA GTGATGTTGG AATCGTTTCT TTTGTTT.GT
 N26910 CTCTTTT.CT CATCACAGAA GTGATGTTGG AATCGTTTCT TTTGTTT.GT
 H16757 CTCTTTTACT CATCACAGAA GTGATGTTGG AATCGTTTCT TTTGTTT.GT
 N27732 CTCTTTT.CT CATCACAGAA GTGATGTTGG AATCGTTTCT TTTGTTT.GT
         1551
                                                         1600
BIKUNIN CTGATTTATG G TTTTTTT AAGTATAAC AAAAGTTTTT TATTAGCATT
 H39840 CTGATTTATG GGTTTTTTTT AAGTAT
 H95233 CTGATTTATG G..TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 H39841 CTGATTTATG G..TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 N30199 CTGATTTATG G..TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 T52966 CIGATITATG G..ITTITIT AAGTATAAAC AAAAGTITIT TATTAGCATT
 N29508 CTGATTTATG G..TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 N26919 CTGATTTATG G..TTTTTTT AAGTNTAAAC AAAAGTTTTT TATTAGCATT
 N26910 CTGATTTATG G..TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 H16757 CTGATTTATG G.. TTTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
 N27732 CTGATTTATG G...TTTTTT AAGTATAAAC AAAAGTTTTT TATTAGCATT
        1601
Bikunin CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
 H95233 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAA
 H39841 CTGAAAGAAG GAAAGTAAAN TGTACAAGTT TAATAAAAAG GGGCCTTCCC
 N30199 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
 T52966 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
 N29508 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
 N26919 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
N26910 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
H16757 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
N27732 CTGAAAGAAG GAAAGTAAAA TGTACAAGTT TAATAAAAAG GGGCCTTCCC
        1651
Bikunin CTTTAG AAT AAAAAAAAA AAAAAAAAA AAAAAAAAA
H39841 CTTTAA.
N30199 CTTTAG. AAT AAA
T52966 CTTTAGGAAT NAAAANAAAA AAGGGTG
N29508 CTTTAG.AAT AAATTTCAGC ATGTGCTTTC AA
N26919 CTTTAG.AAT AAAAAAAAA AAAAAAAAA A
N26910 CTTTAG.AAT AAATTTCAGC ATGTGCTTTC AAAAAA
H16757 CTTTAG.AAT AAAAAAAAA AAAAAAAAA AAAAAA
N27732 CTTTAG. AAT AAAAAAAAA AAAAAAAAA AAAAAAAAA
```

## FIGURE 4D

EST	consens	MLRAEADGVS	RLLGSLLLSG	VLAADRERSI	HDECLVSKVV	GRCRASMPRW	50
EST	consens	WYNVTDGSCQ	LFVYGGCDGN	SNNYLTKEEC	LKKCATVTEN	ATGDLATSRN	100
EST	consens	AADSSVPSAP	RRQDSEDHSS	DMFNYEEYCT	ANAVTGPCRA	SFPRWYFDVE	150
EST	consens	RNSCNNFIYG	GCRGNKNSYR	SEEACMLRCF	RQQENPPLPL	GSK <u>VVVLAGL</u>	200
EST	consens	FVMVLILFLG	ASMVYLIRVA	RRNQERALRT	VWSSGDDKEO	LVKNTYVL	248

# FIGURE 4E

cDNA translation					T	
cDNA	TGATCGCGAG	ACCCCAACGG	CTGGTGGCGT	CGCCTGCGCG	TCTCGGCTGA	53
translation	. S R D	P N G	W W R	R L R V	S A E	-30
cDNA	GCTGGCC <b>ATG</b>	GCGCAGCTGT	GCGGGCTGAG	GCGGAGCCGG	GCGTTTCTCG	10
translation	L A M	A Q L C	G L R	R S R	A F L A	1-13
cDNA	CCCTGCTGGG	ATCGCTGCTC	CTCTCTGGGG	TCCTGGCGGC	CGACCGAGAA	153
translation	L L G	S L L	L S G V	L A A	D R E	4
cDNA	CGCAGCATCC	ACGACTTCTG	CCTGGTGTCG	AAGGTGGTGG	GCAGATGCCG	203
translation	R S I H	D F C	L V S	K V V G	R C R	
cDNA	GGCCTCCATG	CCTAGGTGGT	GGTACAATGT	CACTGACGGA	TCCTGCCAGC	253
translation	A S M	P R W W	Y N V	T D G	S C Q L	38
cDNA	TGTTTGTGTA	TGGGGGCTGT	GACGGAAACA	GCAATAATTA	CCTGACCAAG	303
translation	F V Y	G G C	D G N S	N N Y	L T K	54
cDNA	GAGGAGTGCC	TCAAGAAATG	TGCCACTGTC	ACAGAGAATG	CCACGGGTGA	353
translation	E E C L	K K C	A T V	T E N A	T G D	71
cD <b>NA</b>	CCTGGCCACC	AGCAGGAATG	CAGCGGATTC	CTCTGTCCCA	AGTGCTCCCA	403
translation	L A T	S R N A	A D S	S V P	S A P R	88
cDNA	GAAGGCAGGA	TTCTGAAGAC	CACTCCAGCG	ATATGTTCAA	CTATGAAGAA	453
translation	R Q D	S E D	H S S D	M F N	Y E E	104
cDNA	TACTGCACCG	CCAACGCAGT	CACTGGGCCT	TGCCGTGCAT	CCTTCCCACG	503
translation	Y C T A	N A V	T G P	C R A S	F P R	121
cDNA	CTGGTACTTT	GACGTGGAGA	GGAACTCCTG	CAATAACTTC	ATCTATGGAG	553
translation	W Y F	D V E R	N S C	N N F	I Y G G	138
cDNA	GCTGCCGGGG	CAATAAGAAC	AGCTACCGCT	CTGAGGAGGC	CTGCATGCTC	603
translation	C R G	N K N	S Y R S	E E A	C M L	154
cDNA	CGCTGCTTCC	GCCAGCAGGA	GAATCCTCCC	CTGCCCCTTG	GCTCAAAGGT	653
translation	R C F R	Q Q E	N P P	L P L G	S K <u>Y</u>	171
cDNA	GGTGGTTCTG	GCGGGGCTGT	TCGTGATGGT	GTTGATCCTC	TTCCTGGGAG	703
translation	V V L	A G L F		L L L	F L G A	188
cDNA	CCTCCATGGT	CTACCTGATC	CGGGTGGCAC	GGAGGAACCA	GGAGCGTGCC	753
translation		Y L I	R V A R	R N Q	E R A	204
cDNA translation	CTGCGCACCG L R T V	TCTGGAGCTT W S F	CGGAGATGA G D			782 213

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### FIGURE 4F

CDNA	GCACGAGTTG	GGAGGTGTAG	CGCGGCTCTG	AACGCGCTGA	GGGCCGTTGA	5.0
CDNA	GTGTCGCAGG	CGGCGAGGGC	GCGAGTGAGG	AGCAGACCCA	GGCATCGCGC	100
cDNA	GCCGAGAAGG	CCGGGCGTCC	CCACACTGAA	GGTCCGGAAA	GGCGACTTCC	150
CDNA	GGGGGCTTTG	GCACCTGGCG	GACCCTCCCG	GAGCGTCGGC	ACCTGAACGC	200
CDNA	GAGGCGCTCC	ATTGCGCGTG	CGCGTTGAGG	GGCTTCCCGC	ACCTGATOGO	250
CDNA	GAGACCCCAA	CGGCTGGTGG	CGTCGCCTGC	GCGTCTCGGC	TGAGCTGGCC	300
CDNA	ATGGCGCAGC	TGTGCGGGCT	GAGGCGGAGC	CGGGCGTTTC	TOGGOOTGOT	350
translation	M A Q L	C G L	R R S	RAFL	A L L	-11
cDNA	GGGATCGCTG	CTCCTCTCTG	GGGTCCTGGC	GGCCGACCGA	GAACGCAGCA	400
translation	G S L	L L S G	V L A	A D R	E R S I	-
cDNA	TCCACGACTT	CTGCCTGGTG	TCGAAGGTGG	TGGGCAGATG	CCGGGCCTCC	
translation	H D F	c r A	s k v v	GRC	R A S	23
cDNA	ATGCCTAGGT	GGTGGTACAA	TGTCACTGAC	GGATCCTGCC	AGCTGTTTGT	500
translation	M P R W	W Y N	VTD	G S C Q	L F V	40
CDNA	GTATGGGGGC	TGTGACGGAA	ACAGCAATAA	TTACCTGACC	AACCACCACT	225
translation	Y G G	C D G N	S N N	Y L T	K E E C	55.
					2 2 0	٠, ر
CDNA	GCCTCAAGAA	ATGTGCCACT	GTCACAGAGA	ATGCCACGGG	TGACCTGGCC	600
		CAT				
cDNA	ACCAGCAGGA	ATGCAGCGGA	TTCCTCTGTC	CCAAGTGCTC	CCAGAAGGCA	
translation	TSRN	A A D	s s v	P S A P	RRQ	90
		GACCACTCCA				
translation	D S E	D H S S	D M F	NYE	EYCI	107
- 0.11	6666611666	1656165666		21.00000000		
CDNA		V T G				
translation	ANA	V . 3	PCRA	5 F P	R W Y	123
cDNA	TTTCACCTCC	3686633676	CTCCSSTSSC	TTC\ TCT\ TC	63.666maaaa	222
		R N S				
Clanstacion		V 14 2	C N N	r i i G	GCR	140
CDNA	GGGCAATAAG	AACACTACC	GCTCTGAGGA	GGCCTGCATG	CTCCCCTCCT	050
translation	G N K	N S Y R	S E E	A C M	T B C F	157
c	0 11 11	0	5 5 <u>5</u>	A C II	u R C :	- 5 /
CDNA	TCCGCCAGCA	GGAGAATCCT	CCCCTGCCCC	TTGGCTCAAA	GGTGGTGGTT	900
translation						
	4	•		J J 1.	<del></del>	<b>*</b> * <b>*</b>
CDNA	CTGGCGGGGC	TGTTCGTGAT	GGTGTTGATC	CTCTTCCTGG	GAGCCTCCAT	950
translation						
						. , ,
CDNA	GGTCTACCTG	ATCCGGGTGG	CACGGAGGAA	CCAGGAGCGT	GCCCTGCGCA	1 303
translation						
11331401011				4 - ''	~ .	2 .

...AA TÜÜTTÜRE LEETTI . MARASAB TEESTAA AGGA AGGAAL... translation V & \* 225

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# FIGURE 4F (Con't)

CDNA	ATGTGTGAGC	TTTTTTTAAA	TAGAGGGATT	GACTCGGATT	TGAGTGATCA	1150
CDNA	TTAGGGCTGA	GGTCTGTTTC	TCTGGGAGGT	AGGACGGCTG	CTTCCTGGTC	1200
CDNA	TGGCAGGGAT	GGGTTTGCTT	TGGAAATCCT	CTAGGAGGCT	CCTCCTCGCA	1250
CDNA	TGGCCTGCAG	TCTGGCAGCA	GCCCCGAGTT	GTTTCCTCGC	TGATCGATTT	1300
CDNA	CTTTCCTCCA	GGTAGAGTTT	TCTTTGCTTA	TGTTGAATTC	CATTGCCTCC	1350
CDNA	TTTTCTCNAT	CACAGAAGTG	ATGTTGGAAT	CGTTTCTTTT	GTTTGTCTGA	1400
CDNA	TTTATGGTTT	TTTTAAGTAT	AAACAAAAGT	TTTTTATTAG	CATTCTGAAA	1450
CDNA	GAAGGAAAGT	AAAATGTACA	AGTTTAATAA	AAAGGGGCCT	TCCCCTTTAG	1500
CDNA	AATAAATTTC	CAGCATGTTG	CTTTCAAAAA	$\mathbf{A}\mathbf{A}\mathbf{A}\mathbf{A}\mathbf{A}\mathbf{A}\mathbf{A}\mathbf{A}\mathbf{A}\mathbf{A}$	AAAA	
1550						

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## FIGURE 4G

EST consens			MLR	AEADGVSRLL	GSLLLSGVLA	-1
PCR clone			MAQLCGL	RRSRAFLALL	GSLLLSGVLA	- 1
λcDNA clone			MAQLCGL	RRSRAFLALL	GSLLLSGVLA GSLLLSGVLA GSLLLSGVLA	- 1
EST consens	ADRERSIHDE	CLVSKVVGRC	RASMPRWWYN	VTDGSCQLFV	YGGCDGNSNN	50
PCR clone				_		
AcDNA clone						
EST consens	YLTKEECLKK	CATVTENATG	DLATSRNAAD	SSVPSAPRRO	DSEDHSSDMF	100
PCR clone						
λcDNA clone				_		
EST consens	NYEEYCTANA	VTGPCRASEP	RWYFDVERNS	CNNFIYGGCR	GNKNSYRSEE	150
PCR clone						
λcDNA clone						
EST consens	ACMLRCFRQQ	ENPPLPLGSK	VVVLAGLEVM	VLILELGASM	<u>VYLI</u> RVARRN	200
PCR clone						
λcDNA clone						
<b>505</b>		CCDOVECTUV	MENTAL			225
EST consens			MITAT			
PCR clone						213
λcDNA clone	QERALRTVWS	SGDDKEQLVK	NTYVL			225

% Protease Inhibition

# Purification of Placental Bikunin using Superdex 75 Gel-Filtration

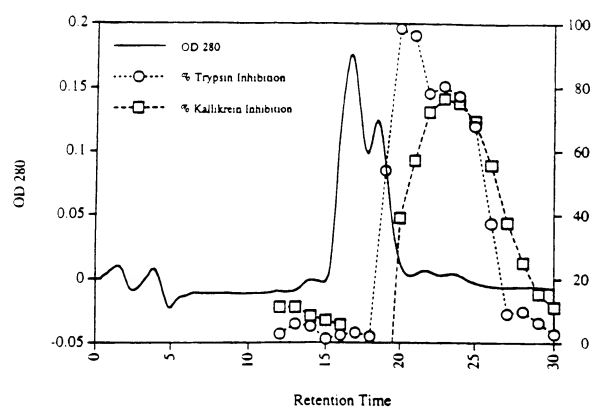
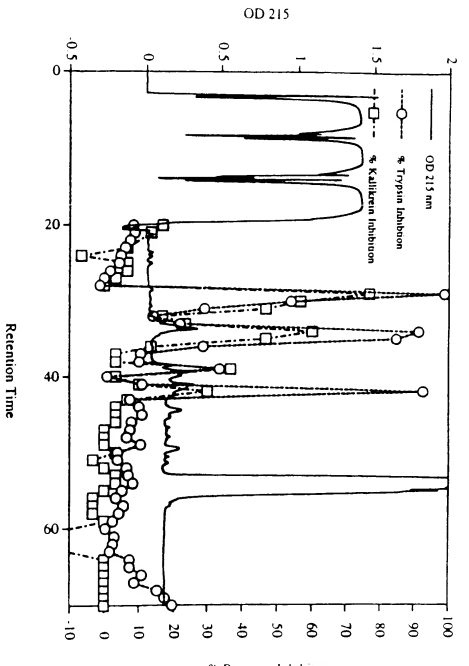


FIGURE 5



FIGURE 6



% Protease Inhibition

Figure 7

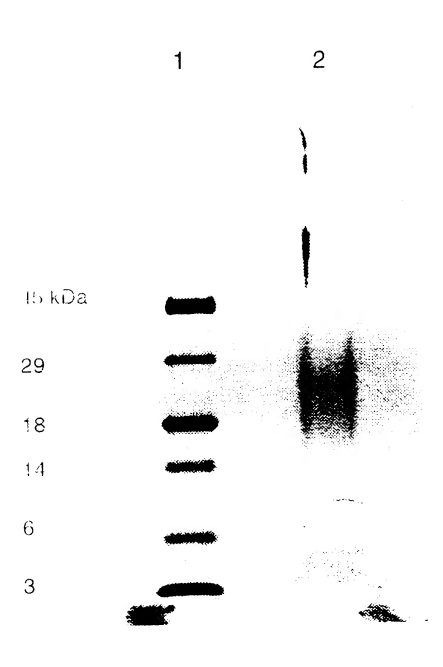


Figure 8A

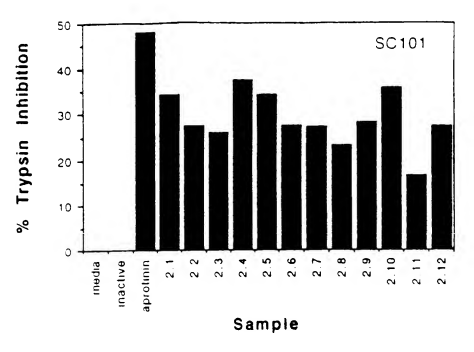
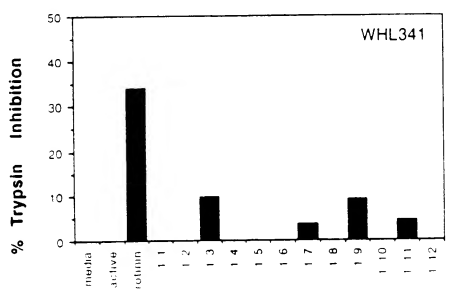
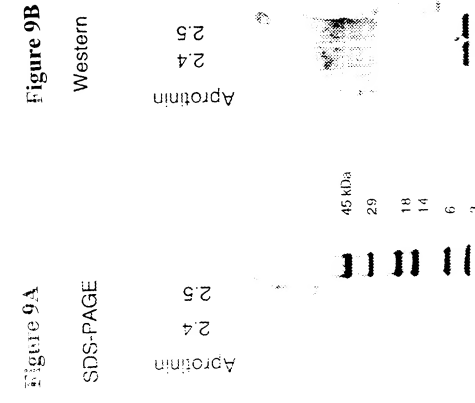


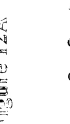
Figure 8B



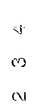


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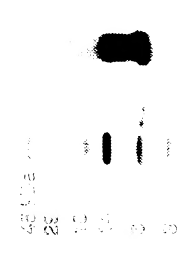
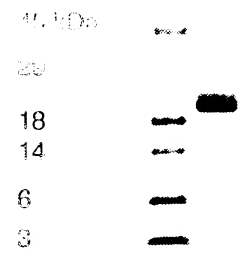




Figure 13

1 2



# Figure 10

1 2

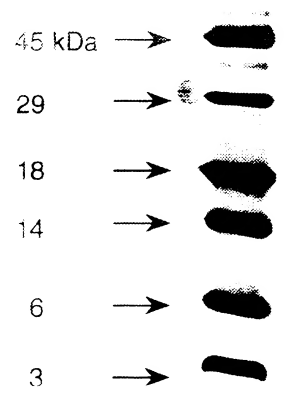


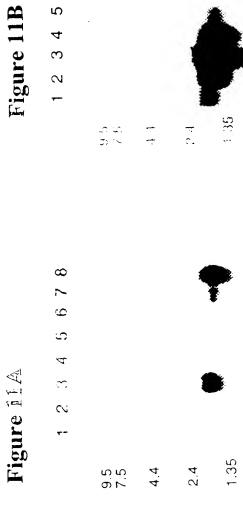
Figure 11A

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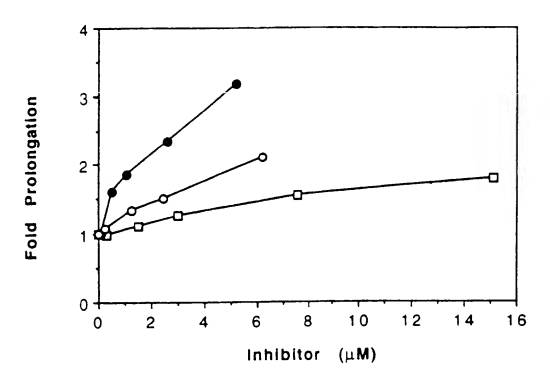
Heart

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Figure 14



Inten 3al Application No PCT/US 97/03894

A. CLASSIFICATION OF SUBJECT MATTER
1PC 6 C12N15/15 C07K14/81 A61K38/57

According to International Patent Classification (IPC) or to both national classification and IPC

#### B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)  $IPC=6=-C07\,K$ 

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	EMBL/GENBANK DATABASES Accession no R35464 Sequence reference HS46499, May 4, 1995 L.HILLIER ET AL: "The WashU-Merck EST Project" XP002039653 see the whole document	1-6,11
X	EMBL/GENBANK DATABASES Accession no N39798 Sequence reference HS798277, January 26, 1996 L. HILLIER ET AL: "The WasU-Merck EST Project" XP002039654 see the whole document	1-6,11

X Further documents are listed in the continuation of box C.	X Patent family members are listed in annex.
* Special categories of cited documents:  A' document defining the general state of the art which is not considered to be of particular relevance.  E' earlier document but published on or after the international filing date.  L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified).  O' document referring to an oral disclosure, use, exhibition or other means.  P' document published prior to the international filing date but later than the priority date claimed.	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention.  "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone.  "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.  "&" document member of the same patent family
Date of the actual completion of the international search 4 September 1997	Date of mailing of the international search report  0 1, 10, 97
Name and mailing address of the ISA	Authorized officer

European Patent Office, P.B. 5818 Patendaan 2 NL - 2280 HV Rijswijk

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Intern at Application No PCT/US 97/03894

Category Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No.			
alegory "	Cuation of document, with indication, where appropriate, of the relevant passages	Kelevani as challi 149	
X	EMBL/GENBANK DATABASES Accession no R74593 Sequence reference HS593137, June 9, 1995 L.HILLIER ET AL: "The WashU-Merck EST Project" XP002039655 see the whole document	1-6,11	
Ρ,Χ	EP 0 758 682 A (MITSUBISHI CHEM CORP) 19 February 1997 see the whole document	1-11	
P,A	JOURNAL OF BIOLOGICAL CHEMISTRY 272 (10). 1997. 6370-6376. ISSN: 0021-9258, XP002039700 SHIMOMURA T ET AL: "Hepatocyte growth factor activator inhibitor, a novel Kunitz-type serine protease inhibitor."		
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A	JOURNAL OF BIOLOGICAL CHEMISTRY 271 (7). 1996. 3615-3618. ISSN: 0021-9258, XP002039701 MIYAZAWA K ET AL: "Activation of hepatocyte growth factor in the injured tissues is mediated by hepatocyte growth factor activator."	7-10	

Intrinational application No

PCT/US 97/03894

Box	Observations where certain claims were found unsearchable (Continuation of Rem 1 of first sneet)
Thus inte	mational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons
1 X	Claims Nos. because they relate to subject matter not required to be searched by this Authority, namely  Remark: Although claim(s) 7-9  is(are) directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.
2	Claims Nos because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful international Search can be carried out, specifically
3	Claims Nos. because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a)
Box II	Observations where unity of invention is tacking (Continuation of item 2 of first sheet)
This Inte	emational Searching Authority found multiple inventions in this international application, as follows
1 🔲	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims
2	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee
3	As only some of the required additional search fees were timely paid by the applicant, this international Search Report covers only those claims for which fees were paid, specifically claims Nos
4	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims, it is covered by claims Nos.
Remar	The additional search fees were accompanied by the applicant's protest  No protest accompanied the payment of additional search fees

unformation on patent family members

Interr 321 Application No
PCT/US 97/03894

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